

IMMUNOGENIC COMPOSITIONS FOR *STREPTOCOCCUS AGALACTIAE*

This application claims the benefit and incorporates by reference in its entirety U.S. provisional application 60/548,789, filed February 26, 2004 and claims priority to International Patent Application No. PCT/US03/29167, Attorney Reference No. PP19766.002, filed on September 15, 2003, incorporated herein in its entirety.

FIELD OF THE INVENTION

The invention relates to an immunogenic antigen derived from *Streptococcus agalactiae* (“GBS”) and its use in combinations with other GBS antigens to provide for broader coverage among different GBS strains. In particular, the invention relates to a composition comprising a combination of two or more GBS antigens, wherein the combination includes GBS 80 or a fragment thereof. The combination may include GBS 80 and at least one other GBS antigen. For example, the combination may include GBS 80 and up to thirteen GBS antigens. In a preferred embodiment, the combination may include GBS 80 and up to ten GBS antigens. In a more preferred embodiment, the combination may include GBS 80 and up to five GBS antigens. In one embodiment, the combination may consist of two to thirteen GBS antigens selected from an antigen group consisting of GBS 80, GBS 91, GBS 104, GBS 184, GBS 276, GBS 305, GBS 322, GBS 330, GBS 338, GBS 361, GBS 404, GBS 690, and GBS 691. Preferably, the combination includes GBS 80 in combination with one or more of GBS 104 and GBS 322.

BACKGROUND OF THE INVENTION

GBS has emerged in the last 20 years as the major cause of neonatal sepsis and meningitis that affect 0.5 – 3 per 1000 live births, and an important cause of morbidity among the older age group affecting 5 – 8 per 100,000 of the population. Current disease management strategies rely on intrapartum antibiotics and neonatal monitoring which have reduced neonatal case mortality from >50% in the 1970’s to less than 10% in the 1990’s. Nevertheless, there is still considerable morbidity and mortality and the management is expensive. 15 – 35% of pregnant women are asymptomatic carriers and at high risk of transmitting the disease to their babies. Risk of neonatal infection is associated with low serotype specific maternal antibodies and high titers are believed to be protective. In addition, invasive GBS disease is increasingly recognized in elderly adults with underlying disease such as diabetes and cancer.

The “B” in “GBS” refers to the Lancefield classification, which is based on the antigenicity of a carbohydrate which is soluble in dilute acid and called the C carbohydrate. Lancefield identified 13 types of C carbohydrate, designated A to O, that could be serologically differentiated. The organisms

that most commonly infect humans are found in groups A, B, D, and G. Within group B, strains can be divided into at least 9 serotypes (Ia, Ib, Ia/c, II, III, IV, V, VI, VII and VIII) based on the structure of their polysaccharide capsule. In the past, serotypes Ia, Ib, II, and III were equally prevalent in normal vaginal carriage and early onset sepsis in newborns. Type V GBS has emerged as an
5 important cause of GBS infection in the USA, however, and strains of types VI and VIII have become prevalent among Japanese women.

The genome sequence of a serotype V strain 2603 V/R has been published (Ref. 1) and various polypeptides for use as vaccine antigens have been identified (Ref. 2). The vaccines currently in clinical trials, however, are based on polysaccharide antigens. These suffer from serotype-
10 specificity and poor immunogenicity, and so there is a need for effective vaccines against *S. agalactiae* infection.

It is an object of the invention to provide further and improved compositions for providing immunity against GBS disease and/or infection. The compositions are based on a combination of two or more (e.g., three or more) GBS antigens.
15

SUMMARY OF THE INVENTION

Applicants have discovered that an immunogenic GBS antigen, GBS 80, is particularly suitable for immunization purposes, especially when used in combination with other GBS antigens. The combination may include GBS 80 and at least one other GBS antigen or up to thirteen other GBS
20 antigens. In a preferred embodiment, the combination may include GBS 80 and up to 10 GBS antigens. In a more preferred embodiment, the combination includes GBS 80 and up to five GBS antigens. In particular, the invention relates to a composition comprising a combination of two or more GBS antigens, wherein the combination includes GBS 80 or a fragment thereof. In one embodiment, the combination may consist of two to thirteen GBS antigens selected from the group
25 consisting of GBS 80, GBS 91, GBS 104, GBS 184, GBS 276, GBS 305, GBS 322, GBS 330, GBS 338, GBS 361, GBS 404, GBS 690, and GBS 691. Preferably, the combination consists of GBS 80, GBS 104 and GBS 322.

Instead of the full length antigen, the combination may comprise an immunogenic fragment of the selected GBS antigen and/or a polypeptide sequence having sequence identity to the selected
30 antigen.

Preferably, the combination of GBS antigens consists of three, four, five, six, seven, eight, nine, or ten GBS antigens. Still more preferably, the combination of GBS antigens consists of three, four, or five GBS antigens.

DETAILED DESCRIPTION OF THE INVENTION

The practice of the present invention will employ, unless otherwise indicated, conventional methods of chemistry, biochemistry, molecular biology, immunology and pharmacology, within the skill of the art. Such techniques are explained fully in the literature. See, e.g., *Remington's Pharmaceutical Sciences*, Mack Publishing Company, Easton, Pa., 19th Edition (1995); *Methods In Enzymology* (S. Colowick and N. Kaplan, eds., Academic Press, Inc.); and *Handbook of Experimental Immunology*, Vols. I-IV (D.M. Weir and C.C. Blackwell, eds., 1986, Blackwell Scientific Publications); Sambrook, et al., *Molecular Cloning: A Laboratory Manual* (2nd Edition, 1989); *Handbook of Surface and Colloidal Chemistry* (Birdi, K.S. ed., CRC Press, 1997); *Short Protocols in Molecular Biology*, 4th ed. (Ausubel et al. eds., 1999, John Wiley & Sons); *Molecular Biology Techniques: An Intensive Laboratory Course*, (Ream et al., eds., 1998, Academic Press); *PCR (Introduction to Biotechniques Series)*, 2nd ed. (Newton & Graham eds., 1997, Springer Verlag); Peters and Dalrymple, *Fields Virology* (2d ed), Fields et al. (eds.), B.N. Raven Press, New York, NY.

All publications, patents and patent applications cited herein, are hereby incorporated by reference in their entireties.

GBS Antigens

5 As discussed above, the invention provides an immunogenic composition comprising a combination of two or more GBS antigens, wherein said combination includes GBS 80 or a fragment thereof.

10 The combinations of GBS antigens may include polypeptide fragments of the identified GBS antigens. The length of the fragment may vary depending on the amino acid sequence of the specific GBS antigen, but the fragment is preferably at least 7 consecutive amino acids, (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200 or more). Preferably the fragment comprises one or more epitopes from the sequence. Other preferred fragments include (1) the N-terminal signal peptides of each identified GBS antigen, (2) the identified GBS antigens without their N-terminal signal peptides, and (3) each identified GBS antigen wherein up to 10 amino acid residues (e.g. 1, 2, 15 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) are deleted from the N-terminus and/or the C-terminus e.g. the N-terminal amino acid residue may be deleted. Other fragments omit one or more domains of the protein (e.g. omission of a signal peptide, of a cytoplasmic domain, of a transmembrane domain, or of an extracellular domain).

20 The combinations of GBS antigens may include polypeptide sequences having sequence identity to the identified GBS antigens. The degree of sequence identity may vary depending on the amino acid sequence (a) in question, but is preferably greater than 50% (e.g. 60%, 65%, 70%, 75%,

80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, 99.5% or more). Polypeptides having sequence identity include homologs, orthologs, allelic variants and functional mutants of the identified GBS antigens. Typically, 50% identity or more between two proteins is considered to be an indication of functional equivalence. Identity between proteins is preferably determined by the

5 Smith-Waterman homology search algorithm as implemented in the MPSRCH program (Oxford Molecular), using an affinity gap search with parameters *gap open penalty*=12 and *gap extension penalty*=1.

The polypeptides can, of course, be prepared by various means (e.g. recombinant expression, purification from GBS, chemical synthesis etc.) and in various forms (e.g. native, fusions, 10 glycosylated, non-glycosylated etc.). They are preferably prepared in substantially pure form (i.e. substantially free from other streptococcal or host cell proteins) or substantially isolated form.

GBS 80

As discussed above, the invention relates to the use of GBS 80 in synergistic combination with other GBS antigens. GBS 80 refers to a putative cell wall surface anchor family protein.

15 Nucleotide and amino acid sequence of GBS 80 sequenced from serotype V isolated strain 2603 V/R are set forth in Ref. 2 as SEQ ID 8779 and SEQ ID 8780. These sequences are also set forth below as SEQ ID NOS 1 and 2:

SEQ ID NO. 1

20 ATGAAATTATCGAAGAAGTTATTGTTTCGGCTGCTGTTAACAAATGGTGGCGGGGTCAACTGTTGA
ACCAGTAGCTCAGTTGCGACTGGAATGAGTATTGTAAGAGCTGCAGAAGTGTACAAGAACGCCAG
CGAAAACAAACAGTAAATATCTATAAATTACAAGCTGATAGTTATAAATCGGAAATTACTCTAATGGT
GGTATCGAGAATAAAGACGGCGAAGTAATATCTAATCTGCTAAACTTGGTACAATGTAAGGTTT
GCAAGGTGTACAGTTAACGTTATAAAGTCAAGACGGATATTCTGTTGATGAAATTGAAAAATTGA
25 CAACAGTTGAAGCAGCAGATGCAAAAGTTGGAACGATTCTGAAAGAAGGTGTCACTCTACCTCAAAAA
ACTAATGCTCAAGGTTGGTCGATGCTCTGGATTCAAAAAGTAATGTGAGATACTTGTATGTAGA
AGATTAAAGAATTACACCTCAAACATTACCAAAGCTTATGCTGTACCGTTGTGTTGGAATTACCAAG
TTGCTAACTCTACAGGTACAGGTTCCCTTCTGAAATTAAATATTACCTAAAAACGTTGAACTGAT
GAACCAAAACAGATAAAGATGTTAAAAAATTAGGTCAAGGACGATGCAAGGTTATACGATTGGTGAAGA
30 ATTCAAATGTTCTGAAATCTACAATCCCTGCCAATTAGGTGACTATGAAAAATTGAAATTACTG
ATAAAATTGCGATGGCTTGACTTATAAATCTGTTGGAAAATCAAGATTGGTTCGAAAACACTGAAT
AGAGATGAGCACTACACTATTGATGAACCAACAGTTGATAACCAAAATACATTAAAATACGTTAA
ACCAGAGAAATTAAAGAAATTGCTGAGCTACTTAAAGGAATGACCCCTGTTAAAATCAAGATGCTC
35 TTGATAAAGCTACTGCAAATACAGATGATGCGGCATTGGAAATTCCAGTTGCAACTATTAAAT
GAAAAAGCAGTTAGGAAAAGCAATTGAAAATACTTGAACCTCAATATGACCATACCTCCTGATAA
AGCTGACAATCCAAAACCATCTAATCCTCAAGAAAACCAGAAGTTCATACTGGTGGGAAACGATTG
TAAAGAAAAGACTCAACAGAAACACAAACACTAGGTGGTGCTGAGTTGATTGTTGGCTCTGATGGG
ACAGCAGTAAATGGACAGATGCTCTTATAAAGCAATACTAATAAAAACATATTGCTGGAGAAC
40 TGTTACTGGCAACCAATCAAATTGAAATCACATACAGACGGTACGTTGAGATTAAGGTTGGCTT
ATGCAGTTGATGCGAATGCAGAGGGTACAGCAGTAACCTACAAATTAAAAGAAACAAAGCACCAGAA
GGTTATGTAATCCCTGATAAAGAAATCGAGTTACAGTATCACAAACATCTTATAATACAAAACCAAC
TGACATCACGGTTGATAGTGCTGATGCAACACACTGATACAATTAAAACAACAAACGTCCTCAATCC
CTAATACTGGTGGTATTGGTACGGCTATCTTGTGCTATCGGTGCTGCGGTGATGGCTTTGCTGTT
AAGGGGATGAAGCGTCGTACAAAGATAAC

SEQ ID NO: 2

5 MKLSKKLLFSAAVLTMAGSTVEPVAQFATGMSIVRAAEVSQERPAKTTVNIYKLQADSYKSEITSNG
 GIENKDGEVISNYAKLGDNVKGLQGVQFKRYKVKTDISVDELKKLTVEAADAKVGTILEEGVSLPQK
 TNAQGLVVDALDSKSNVRYLYVEDLKNSPSNITKAYAVPFVLELPVANSTGTGFLSEINIYPKNVVTD
 EPKTDKDVKKLGQDDAGYTIGEEFKWFLKSTIPANLGDYEKFEITDKFADGLTYKSVGKIKIGSKTLN
 RDEHYTIDEPVDNQNTLKITFKPEKFKEIAELLKGMTLVKNQDALKATANTDDAAFLEIPVASTIN
 10 EKAVLGKAIENTFELQYDHTPDKADNPKPSNPPRKPEVHTGGKRFVKKDSTETQTLGGAEFDLLASDG
 TAVKWTDALIKANTNKNYIAGEAVTGQPIKLKSHTDGTFEIKGLAYAVDANAEGTAVTYKLKETKAP
 GYVIPDKEIEFTVSQTSYNTKPTDITVDSADATPDTIKNNKRPSIPNTGGIGTAIFVAIGAAVMAFAV
 KGMKRRTKDN

15 As described above, the combinations of the invention may include a fragment of a GBS antigen. In some instances, removal of one or more domains, such as a leader or signal sequence region, a transmembrane region, a cytoplasmic region or a cell wall anchoring motif, may facilitate cloning of the gene encoding the antigen and/or recombinant expression of the GBS protein. In addition, fragments comprising immunogenic epitopes of the cited GBS antigens may be used in the compositions of the invention.

20 GBS 80 contains an N-terminal leader or signal sequence region which is indicated by the underlined sequence at the beginning of SEQ ID NO: 2 above. In one embodiment, one or more amino acids from the leader or signal sequence region of GBS 80 are removed. An example of such a GBS 80 fragment is set forth below as SEQ ID NO: 3:

SEQ ID NO: 3

25 AEVSQERPAKTTVNIYKLQADSYKSEITSNGGIENKDGEVISNYAKLGDNVKGLQGVQFKRYKVKTDI
 SVDELKKLTVEAADAKVGTILEEGVSLPQKTNAQGLVVDALDSKSNVRYLYVEDLKNSPSNITKAYA
 VPVLELPVANSTGTGFLSEINIYPKNVVTDEPKTDKDVKKLGQDDAGYTIGEEFKWFLKSTIPANLG
 30 DYEKFEITDKFADGLTYKSVGKIKIGSKTLNRDEHYTIDEPVDNQNTLKITFKPEKFKEIAELLKGM
 TLVKNQDALKATANTDDAAFLEIPVASTINEKAVLGKAIENTFELQYDHTPDKADNPKPSNPPRKPE
 VHTGGKRFVKKDSTETQTLGGAEFDLLASDGTAVKWTDALIKANTNKNYIAGEAVTGQPIKLKSHTD
 GTFEIKGLAYAVDANAEGTAVTYKLKETKAPEGYVIPDKEIEFTVSQTSYNTKPTDITVDSADATPDTI
 KNNKRPSIPNTGGIGTAIFVAIGAAVMAFAVKGMKRRTKDN

35 GBS 80 contains a C-terminal transmembrane region which is indicated by the underlined sequence near the end of SEQ ID NO: 2 above. In one embodiment, one or more amino acids from the transmembrane region and/or a cytoplasmic region are removed. An example of such a GBS 80 fragment is set forth below as SEQ ID NO: 4:

SEQ ID NO: 4

40 MKLSKKLLFSAAVLTMAGSTVEPVAQFATGMSIVRAAEVSQERPAKTTVNIYKLQADSYKSEITSNG
 GIENKDGEVISNYAKLGDNVKGLQGVQFKRYKVKTDISVDELKKLTVEAADAKVGTILEEGVSLPQK
 TNAQGLVVDALDSKSNVRYLYVEDLKNSPSNITKAYAVPFVLELPVANSTGTGFLSEINIYPKNVVTD
 EPKTDKDVKKLGQDDAGYTIGEEFKWFLKSTIPANLGDYEKFEITDKFADGLTYKSVGKIKIGSKTLN
 RDEHYTIDEPVDNQNTLKITFKPEKFKEIAELLKGMTLVKNQDALKATANTDDAAFLEIPVASTIN
 45 EKAVLGKAIENTFELQYDHTPDKADNPKPSNPPRKPEVHTGGKRFVKKDSTETQTLGGAEFDLLASDG

TAVKWTDALIKANTNKNYIAGEAVTGQPIKLKSHTDGTFEIKGLAYAVDANAEGTAVTYKLKETKAPE
GYVIPDKEIEFTVSQTSYNTKPTDITVDSADATPDTIKNNKRPS *IPNTG*

GBS 80 contains an amino acid motif indicative of a cell wall anchor: **SEQ ID NO: 5**

5 IPNTG (shown in italics in SEQ ID NO: 2 above). In some recombinant host cell systems, it may be preferable to remove this motif to facilitate secretion of a recombinant GBS 80 protein from the host cell. Accordingly, in one preferred fragment of GBS 80 for use in the invention, the transmembrane and/or cytoplasmic regions and the cell wall anchor motif are removed from GBS 80. An example of such a GBS 80 fragment is set forth below as SEQ ID NO: 6.

10

SEQ ID NO: 6

MKLSKKLLFSAAVLTMVAGSTVEPVAQFATGMSIVRAAEVSQERPAKTTVNIYKLQADSYKSEITSNG
GIENKDGEVISNYAKLGDNVKGLQGVQFKRYKVKTDI SVDELKKLTVEAADAKVGTILEEGVSLPQK
TNAQGLVVDALDSKSNVRYLYVEDLKNSPSNITKAYAVPFVLELPVANSTGTGFLSEINIYPKNVVTD
15 EPKTDKDVKKLGQDDAGYTIGEEFKWFLKSTIPANLGDYEKFEITDKFADGLTYKSVGKIKIGSKTLN
RDEHYTIDEPTVDNQNTLKITFKPEKFEIAELLKGMTLVKNQDALKATANTDDAAFLEIPVASTIN
EKAVLGKAIENTFELQYDHTPDKADNPKPSNPPRKPEVHTGGKRFVKKDSTETQTLGGAEFDLLASDG
20 TAVKWTDALIKANTNKNYIAGEAVTGQPIKLKSHTDGTFEIKGLAYAVDANAEGTAVTYKLKETKAPE
GYVIPDKEIEFTVSQTSYNTKPTDITVDSADATPDTIKNNKRPS

20

Alternatively, in some recombinant host cell systems, it may be preferable to use the cell wall anchor motif to anchor the recombinantly expressed protein to the cell wall. The extracellular domain of the expressed protein may be cleaved during purification or the recombinant protein may be left attached to either inactivated host cells or cell membranes in the final composition.

25

In one embodiment, the leader or signal sequence region, the transmembrane and cytoplasmic regions and the cell wall anchor motif are removed from the GBS 80 sequence. An example of such a GBS 80 fragment is set forth below as SEQ ID NO: 7.

30

SEQ ID NO: 7

AEVSQERPAKTTVNIYKLQADSYKSEITSNGGIENKDGEVISNYAKLGDNVKGLQGVQFKRYKVKTDI
SVDELKKLTVEAADAKVGTILEEGVSLPQKTNQGLVVDALDSKSNVRYLYVEDLKNSPSNITKAYA
VPFVLELPVANSTGTGFLSEINIYPKNVVTDEPKTDKDVKKLGQDDAGYTIGEEFKWFLKSTIPANLG
35 DYEKFEITDKFADGLTYKSVGKIKIGSKTLNRDEHYTIDEPTVDNQNTLKITFKPEKFEIAELLKG
MTLVKNQDALKATANTDDAAFLEIPVASTINEKAVLGKAIENTFELQYDHTPDKADNPKPSNPPRKPE
VHTGGKRFVKKDSTETQTLGGAEFDLLASDGTAVKWTDALIKANTNKNYIAGEAVTGQPIKLKSHTDG
40 TFEIKGLAYAVDANAEGTAVTYKLKETKAPEGYVIPDKEIEFTVSQTSYNTKPTDITVDSADATPDTI
KNNKRPS

40

Applicants have identified a particularly immunogenic fragment of the GBS 80 protein. This immunogenic fragment is located towards the N-terminus of the protein and is underlined in the GBS 80 SEQ ID NO: 2 sequence below. The underlined fragment is set forth below as SEQ ID NO: 8.

SEQ ID NO: 2

5 MKLSKKLLFSAAVLTMVAGSTVEPVAQFATGMSIVRAAEVSQERPAKTTVNIYKLQADSYKSEITSNG
GIENKDGEVISNYAKLGDNVKGLOGVQFKRYKVKTDISVDELKKLTVEAADAKVGTILEEGVSLPQK
TNAQGLVVDALDSKSNVRYLYVEDLKNSPSNITKAYAVPFVLELPVANSTGTGFLSEINIYPKNVTD
EPKTDKDVKKLGQDDAGYTIGEEFKWFLKSTIPANLGDYEKFEITDKFADGLTYKSVGKIKIGSKTLN
RDEHYTIDEPTVDNQNTLKTFKPEKFKEIAELLKGMTLVKNQDALDKATANTDDAAFLEIPVASTIN
10 EKAVLGKAIENTFELQYDHTPDKADNPKPSNPPRKPEVHTGGKRFVKKDSTETQTLGGAEDLLASDG
TAVKWTDALIKANTNKNYIAGEAVTGQPIKLKSHTDGTFEIKGLAYAVDANAEGTAVTYKLKETKAP
GYVIPDKEIEFTVSQTSYNTKPTDITVDSADATPDTIKNNKRPSIPNTGGIGTAIFVAIGAAVMAFAV
KGMKRRTKDN

SEQ ID NO: 8

15 AEVSQERPAKTTVNIYKLQADSYKSEITSNGGIENKDGEVISNYAKLGDNVKGLOGVQFKRYKVKTDI
SVDELKKLTVEAADAKVGTILEEGVSLPQKTNQGLVVDALDSKSNVRYLYVEDLKNSPSNITKAYA
VPFVLELPVANSTGTGFLSEINIYPKNVTDDEPKTDKDVKKLGQDDAGYTIGEEFKWFLKSTIPANLG
DYEKFEITDKFADGLTYKSVGKIKIGSKTLNRDEHYTIDEPTVDNQNTLKTFKPEKFKEIAELLKG

20 The immunogenicity of the protein encoded by SEQ ID NO: 7 was compared against PBS,
GBS whole cell, GBS 80 (full length) and another fragment of GBS 80, located closer to the C-
terminus of the peptide (SEQ ID NO: 9, below).

SEQ ID NO: 9

25 MTLVKNQDALDKATANTDDAAFLEIPVASTINEKAVLGKAIENTFELQYDHTPDKADNPKPSNPPRKPEVHTGGK
RFVKKDSTETQTLGGAEDLLASDGTAVKWTDALIKANTNKNYIAGEAVTGQPIKLKSHTDGTFEIKGLAYAVDA
NAEGTAVTYKLKETKAPEGYVIPDKEIEFTVSQTSYNTKPTDITVDSADATPDTIKNNKRPS

Both an Active Maternal Immunization Assay and a Passive Maternal Immunization Assay
were conducted on this collection of proteins.

30 As used herein, an Active Maternal Immunization assay refers to an *in vivo* protection assay
where female mice are immunized with the test antigen composition. The female mice are then bred
and their pups are challenged with a lethal dose of GBS. Serum titers of the female mice during the
immunization schedule are measured as well as the survival time of the pups after challenge.

35 Specifically, the Active Maternal Immunization assays referred to herein used groups of four
CD-1 female mice (Charles River Laboratories, Calco Italy). These mice were immunized
intraperitoneally with the selected proteins in Freund's adjuvant at days 1, 21 and 35, prior to
breeding. 6-8 weeks old mice received 20 µg protein/dose when immunized with a single
antigen, 30-45 µg protein/dose (15 µg each antigen) when immunized with combination of
antigens. The immune response of the dams was monitored by using serum samples taken on day
0 and 49. The female mice were bred 2-7 days after the last immunization (at approximately t=

40 36 – 37), and typically had a gestation period of 21 days. Within 48 hours of birth, the pups were
challenged via I.P. with GBS in a dose approximately equal to an amount which would be
sufficient to kill 70 – 90 % of unimmunized pups (as determined by empirical data gathered from

PBS control groups). The GBS challenge dose is preferably administered in 50 μ l of THB medium. Preferably, the pup challenge takes place at 56 to 61 days after the first immunization. The challenge inocula were prepared starting from frozen cultures diluted to the appropriate concentration with THB prior to use. Survival of pups was monitored for 5 days after challenge.

5 As used herein, the Passive Maternal Immunization Assay refers to an *in vivo* protection assay where pregnant mice are passively immunized by injecting rabbit immune sera (or control sera) approximately 2 days before delivery. The pups are then challenged with a lethal dose of GBS.

10 Specifically, the Passive Maternal Immunization Assay referred to herein used groups of pregnant CD1 mice which were passively immunized by injecting 1 ml of rabbit immune sera or control sera via I.P., 2 days before delivery. Newborn mice (24-48 hrs after birth) are challenged via I.P. with a 70 - 90% lethal dose of GBS serotype III COH1. The challenge dose, obtained by diluting a frozen mid log phase culture, was administered in 50 μ l of THB medium.

15 For both assays, the number of pups surviving GBS infection was assessed every 12 hrs for 4 days. Statistical significance was estimated by Fisher's exact test.

15 The results of each assay for immunization with SEQ ID NO: 7, SEQ ID NO: 8, PBS and GBS whole cell are set forth in Tables 1 and 2 below.

TABLE 1: Active Maternal Immunization

Antigen	Alive/total	%Survival	Fisher's exact test
PBS (neg control)	13/80	16%	
GBS (whole cell)	54/65	83%	P<0.00000001
GBS80 (intact)	62/70	88%	P<0.00000001
GBS80 (fragment) SEQ ID 7	35/64	55%	P=0.0000013
GBS80 (fragment) SEQ ID 8	13/67	19%	P=0.66

Table 2: Passive Maternal Immunization

Antigen	Alive/total	%Survival	Fisher's exact test
PBS (neg control)	12/42	28%	
GBS (whole cell)	48/52	92%	P<0.00000001
GBS80 (intact)	48/55	87%	P<0.00000001
GBS80 (fragment) SEQ ID 7	45/57	79%	P=0.0000006
GBS80 (fragment) SEQ ID 8	13/54	24%	P=1

20 As shown in Tables 1 and 2, immunization with the SEQ ID NO: 7 GBS 80 fragment provided a substantially improved survival rate for the challenged pups than the comparison SEQ ID NO: 8 GBS 80 fragment. These results indicate that the SEQ ID NO: 7 GBS 80 fragment may comprise an important immunogenic epitope of GBS 80.

Combinations including GBS 80

The invention includes combinations of two or more GBS antigens wherein the combination includes GBS 80 or a fragment thereof. Applicants have discovered that GBS 80 is particularly suitable for immunization in combination with other GBS antigens and that these antigen combinations provide for a broader coverage among different GBS strains.

5 Preferably, the combination of GBS antigens consists of three, four, five, six, seven, eight, nine, or ten GBS antigens. Still more preferably, the combination of GBS antigens consists of three, four, or five GBS antigens.

10 Preferably, the combinations of the invention provide for improved immunogenicity over the immunogenicity of the antigens when administered alone. Improved immunogenicity may be measured, for example, by the Active Maternal Immunization Assay. As discussed above, this assay may be used to measure serum titers of the female mice during the immunization schedule as well as the survival time of the pups after challenge. Preferably, immunization with the immunogenic compositions of the invention yield an increase of at least 2 percentage points (preferably at least 3, 4 15 or 5 percentage points) in the percent survival of the challenged pups as compared to the percent survival from maternal immunization with a single antigen of the composition when administered alone. Preferably, the increase is at least 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29 or 30 percentage points. Preferably, the GBS combinations of the invention comprising GBS 80 demonstrate an increase in the percent survival as compared to the percent 20 survival from immunization with a non-GBS 80 antigen alone.

According to one embodiment of the invention, combinations of antigens or fusion proteins containing a portion or portions of the antigens will include GBS 80 or a portion thereof in combination with from one to 10 antigens, preferably one to 10 or less antigens. Such other antigens include by way of example and not limitation, GBS 67, GBS 91, GBS 104, GBS 184, GBS 276, GBS 25 305, GBS 322, GBS 330, GBS 338, GBS 361, GBS 404, GBS 690, and GBS 691. Still other antigens are identified in U.S. Serial Number 10/415,182, filed April 28, 2003, hereby incorporated in its entirety.

30 Combinations, for example, can include GBS 80, GBS 104, GBS 322, and GBS 276, ; GBS 80, GBS 338, GBS 330; GBS 80, GBS 330, GBS 104; GBS 80, GBS 104, GBS 404; GBS 80, GBS 338, GBS 104; GBS 80, GBS 338, GBS404; GBS 338, GBS 330, GBS 104; GBS 338, GBS 104, GBS 404; GBS 80, GBS 330, GBS 404; GBS 80, GBS 322, GBS 104; GBS 80, GBS 322, GBS 276; GBS 80, GBS 322, GBS 91; GBS 80, GBS 104, GBS 276; GBS 80, GBS 104, GBS 91; GBS 80, GBS 276, GBS 91; GBS 80, GBS 322, GBS 104; GBS 80, GBS 322, GBS 276; GBS 80, GBS 322, GBS 91; GBS 80, GBS 104, GBS 276; GBS 80, GBS 104, GBS 91; GBS 80, GBS 276, GBS 91; GBS 80, GBS 690, GBS 691; GBS 80, GBS 690, GBS 338; GBS 80, GBS 690, GBS 305; GBS 80, GBS 691,

GBS 305; GBS 80, GBS 338, GBS 305; GBS 80, GBS 338, GBS 361; GBS 80, GBS 305, GBS 361; GBS 80, GBS 184, GBS 691; GBS 80, GBS 691, GBS 338; GBS 80, GBS 104, GBS 276, GBS 322; GBS 80, GBS 104, GBS 67, and GBS 322. Examples of combinations of the invention which demonstrate improved immunogenicity are set forth below. A more detailed description of the GBS 5 antigens referred to in these experiments is set forth following the examples.

EXAMPLE 1: Active Maternal Immunization Assay of GBS 80 alone vs. in combination

In this example, the Active Maternal Immunization Assay was used to measure the percent 10 survival of pups challenged with a Type III serotype of GBS (COH1 isolate), at t=56 days. The maternal mice were immunized according to the Active Maternal Immunization Assay schedule discussed above with GBS 80 alone, combinations of GBS antigens (with and without GBS 80), placebo (PBS) or inactivated whole cell GBS isolate as indicated in Table 3 below. In these 15 experiments, the challenge dose for GBS Type III, strain isolate COH1 sufficient to kill 70 – 90 % of unimmunized pups is approximately equal to 10 x LD 50% (where LD 50% is the statistically derived Median Lethal Dose).

Table 3: Active Maternal Immunization Assay of GBS 80 alone vs. in combination

α -GBS	I Challenge t=56 days Type III COH1 10 x LD 50%	
	Alive/treated	Survival %
α -PBS	3/26	11
α -GBS III	9/20	45
80	24/34	70
80+338+330	39/40	97
80+330+104	38/40	95
80+104+404	24/24	100
80+338+104	33/34	97
80+338+404	30/30	100
338+330+104	22/30	73
338+104+404	24/37	65
80+330+404	25/28	89

20 As shown in Table 3, combinations of GBS antigens which included GBS 80 demonstrated an improved immunogenicity over the use of the antigens alone. For example, immunization with GBS 80 alone yielded a 70% survival rate among the challenged pups. Immunization with combinations of GBS 80 with GBS 338, GBS 330, GBS 104, and GBS 404 yielded 95 to 100% survival rate among the challenged pups. This is an increase of 25 to 30 percentage points.

25 By comparison, combinations of these antigens which did not include GBS 80 failed to achieve the % survival of GBS 80 alone. For example, immunization with GBS 338, GBS 104 and

GBS 404 yielded a 65% survival rate. Replacement of any one of these antigens with GBS 80 dramatically increased the percent survival rate to between 97 and 100%. This is an increase of 32 to 35 percentage points. (See percent survival rates of GBS 80, 338, 101 (97%); GBS 80, 338, 404 (100%) and GBS 80, 104, 404 (100%)). Similarly, immunization with GBS 338, 330 and 104 yielded 5 a 73% survival rate. Replacement of any one of these antigens with GBS 80 increased the percent survival rate to between 95 – 97%.

These findings indicate that protection from COH1 isolate is increased with use of GBS 80 in combination with other GBS antigens.

10 **EXAMPLE 2: Active Maternal Immunization Assay of GBS 80, GBS 322, GBS 276, GBS 104 alone vs. in combination**

In this example, the Active Maternal Immunization Assay was used to measure the percent survival of pups challenged with a Type III serotype of GBS (COH1 isolate) at t=56 days. The 15 maternal mice were immunized according to the Active Maternal Immunization Assay schedule discussed above with a single GBS antigen, combinations of GBS antigens with GBS 80, and placebo (PBS) as indicated in Table 4 below.

20 Table 4: Active Maternal Immunization Assay of GBS 80, GBS 322, GBS 276 or GBS 104 alone vs. in combination with GBS 80

α -GBS	I Challenge t=56 days	
	Type III COH1 10x LD 50%	Survival %
Alive/treated		
80 + 322 + 104	27/27	100
80 + 322 + 276	35/38	92
80 + 322 + 91	24/24	100
80 + 104 + 276	29/30	97
80 + 104 + 91	36/40	90
80 + 276 + 91	33/40	82
GBS 80	24/30	80
GBS 322	7/40	17
GBS 276	13/37	35
GBS 104	28/38	74
α -PBS	2/27	7

As shown in Table 4, the combinations of the antigens with GBS 80 yielded improved immunogenicity over the use of the antigens alone. For example, immunization with GBS 322 alone yielded a 17 % survival rate among the challenged pups. Immunization with combinations of GBS 25 322 with GBS 80 and another GBS antigen yielded survival rates of 92 – 100%. As another example, immunization with GBS 104 alone yielded a 74% survival rate. Immunization with combinations of

GBS 104 with GBS 80 and another GBS antigen yielded survival rates of 90 – 100%. As another example, immunization with GBS 276 alone yielded a 35% survival rate. Immunization with combinations of GBS 276 with GBS 80 and another GBS antigen yielded survival rates of 82 – 97%.

Having demonstrated the immunogenicity of the above-described combinations, the duration 5 of the immune response in the mouse model was further analysed. The maternal mice used in the above described Active Maternal Immunization Assay were mated a second time and the resulting pups challenged with a different GBS serotype (Type V, CJB 111 isolate) at a dramatically higher dose (300x LD 50%) at t=91 days. The parameters of this second, much stronger challenge were outside those of the standard Active Maternal Immunization Assay and were meant to probe the limits 10 of the immunological memory generated from the original maternal immunization in the mouse model. Indication of immunological memory in this model under these conditions is thought to be significant. As shown in Table 5, even under these extreme conditions, increased survival rates were generally achieved, particularly for the combination comprising GBS 80, GBS 322 and GBS 104. It was surprising to note that the percent survival rate for the combination of GBS 80, GBS 233 and 15 GBS 104 was 100% for both the first and second challenges.

Table 5: Second generation pups challenged with higher dose of different strain

α-GBS	II Challenge t=91 days Type V CJB111 300x LD 50%	Alive/treated	Survival %
80 + 322 + 104		20/20	100
80 + 322 + 276		32/37	86
80 + 322 + 91		27/30	90
80 + 104 + 276		22/37	59
80 + 104 + 91		36/39	92
80 + 276 + 91		23/28	82
GBS 80		13/30	43
GBS 322		25/30	83
GBS 276		18/40	45
GBS 104		21/39	54
α-PBS		9/36	25

20 **EXAMPLE 3: Active Maternal Immunization Assay of combinations of
GBS 80 with GBS 690, GBS 691, GBS 338, GBS 305, GBS 361 and GBS 184**

In this example additional combinations of GBS antigens were used in the Active Maternal Immunization Assay, again with a GBS Type III COH1 isolate challenge. The maternal mice were immunized according to the Active Maternal Immunization Assay schedule described above with the 25 combinations of GBS antigens set forth in Table 6 below.

Table 6: Active Maternal Immunization Assay using combinations of GBS 80 with GBS 690, GBS 691, GBS 338, GBS 305, GBS 361 and GBS 184

α -GBS	I Challenge t=56 days		
	Type III COH1 10x LD 50%	Alive/treated	Survival %
80 + 690 + 691	26/29	90	
80 + 690 + 338	35/40	87	
80 + 690 + 305	34/35	97	
80 + 691 + 305	37/40	92	
80 + 338 + 305	25/30	83	
80 + 338 + 361	26/30	87	
80 + 305 + 361	23/30	77	
80 + 184 + 691	32/39	82	
α -PBS	10/40	25	

5 The maternal mice in this model were also mated a second time and the resulting pups challenged with the same GBS isolate at a dramatically higher dose (100x LD 50%) at t=84 days. As in the example above, the parameters of this second, much stronger challenge were outside those of the standard Active Maternal Immunization Assay and were meant to probe the limits of the immunological memory generated from the original maternal immunization in the mouse model. As
10 shown in Table 7, even under these extreme conditions, some of the survival rates remained at or above 70%. Surprisingly, the percent survival rates for the combination of GBS 80, GBS 184 and GBS 691 actually increased.

Table 7: Second generation pups challenged with higher dose

α -GBS	II Challenge t=84 days		
	Type III COH1 100x LD 50%	Alive/treated	Survival %
80 + 690 + 691	19/39	49	
80 + 690 + 338	21/30	70	
80 + 690 + 305	23/40	57	
80 + 691 + 305	22/30	73	
80 + 338 + 305	18/30	60	
80 + 338 + 361	25/40	62	
80 + 305 + 361	21/30	70	
80 + 184 + 691	35/40	87	
α -PBS	4/20	20	

EXAMPLE 4: Active Maternal Immunization Assay using combinations of GBS 80 with GBS 690, GBS 691, GBS 338, GBS 305, and GBS 361

In this example additional combinations of GBS antigens were used in the Active Maternal

5 Immunization Assay, this time with a GBS Type V, CJB111 isolate challenge. In these experiments, the challenge dose for the GBS Type V, CJB111 isolate sufficient to kill 70 – 90% of unimmunized pups is approximately equal to $60 \times LD\ 50\%$ (where LD 50% is the statistically derived Median Lethal Dose). The maternal mice were immunized according to the Active Maternal Immunization Assay schedule described above with the combinations of GBS antigens set forth in Table 8 below.

10 As shown in Table 8, in this particular challenge study with this specific Type V strain isolate, the survival rates for all of the combinations achieved at least 70%.

Table 8: Active Maternal Immunization Assay using combinations of GBS 80 with GBS 690, GBS 691, GBS 338, GBS 305 and GBS 361

α -GBS	I Challenge t=56 days		
	Type V CJB111 60x LD 50%	Alive/treated	Survival %
80 + 690 + 691	24/30	80	
80 + 690 + 338	11/17	70	
80 + 691 + 338	7/10	70	
80 + 691 + 305	21/30	70	
80 + 338 + 305	26/30	87	
80 + 338 + 361	26/30	87	
80 + 305 + 361	28/30	93	
GBS 80	21/30	70	
α -PBS	5/18	28	

15 The maternal mice in this model were also mated a second time and the resulting pups challenged with the same GBS isolate at a dramatically higher dose (600x LD 50%) at t=84 days. As in the example above, the parameters of this second, much stronger challenge were outside those of the standard Active Maternal Immunization Assay and were meant to probe the limits of the

20 immunological memory generated from the original maternal immunization in the mouse model. As shown in Table 9, even under these extreme conditions, some of the survival rates remained above 70%. Surprisingly, the percent survival for two of the antigen groups actually increased (GBS 80, GBS 690 and GBS 338) and (GBS 80, GBS 691 and GBS 338).

Table 9: Second generation pups challenged with higher dose

α -GBS	II Challenge t=84 days	
	Type V CJB111 600x LD 50%	Survival %
Alive/treated		
80 + 690 + 691	27/37	73
80 + 690 + 338	15/20	75
80 + 691 + 338	27/30	90
80 + 691 + 305	23/40	57
80 + 338 + 305	12/20	60
80 + 338 + 361	24/30	80
80 + 305 + 361	24/30	80
GBS 80	24/30	80
α -PBS	ND	ND

EXAMPLE 5: Active Maternal Immunization Assay using combinations of GBS 80 with GBS 104, GBS 276, and GBS 322

5 In this example an additional combination of GBS antigens was used in the Active Maternal Immunization Assay, this time with an isolate challenge of different GBS strains. In these experiments, the challenge dose for the different GBS strains was sufficient to kill 60 – 90% of unimmunized pups and is equal to 10 x LD 50% (where LD 50% is the statistically derived Median 10 Lethal Dose). The maternal mice were immunized according to the Active Maternal Immunization Assay schedule described above with the combination of GBS 80 antigen with GBS 104, GBS 276, and GBS 322 antigens in the GBS strains set forth in Table 10 below. Survival % was observed with the GBS combination with two different adjuvants, Alum and Freunds. As shown in Tables 10 and 11, 15 in this particular challenge study, the survival rates for the combination in all of the GBS strains achieved up to 96%.

Table 10: Active Maternal Immunization Assay using combinations of GBS 80 with GBS 104, GBS 276, and GBS 322 – Alum adjuvant

ALUM					
GBS strains	Type	Mix=322+80+104+276		PBS	
		Alive/treated	Survival %	Alive/treated	Survival %
JM9130013	VIII	32/36	89	18/46	40
CJB111	V	118/145	81	21/110	19
COH1	III	96/115	83	22/104	21
M781	III	42/52	81	18/48	38
2603	V	79/145	54	28/128	22
18RS21	II	86/186	46	24/131	18
DK21	II	31/140	22	28/118	24
7357b –	Ib	25/88	28	25/106	23
A909	Ia	4/40	10	9/60	15
090	Ia	2/31	6	4/53	7
SMO53	VII	17/54	31	4/39	10

5 Table 11: Active Maternal Immunization Assay using combinations of GBS 80 with GBS 104, GBS 276, and GBS 322 – Freund adjuvant

Freund					
GBS strains	Type	Mix=322+80+104+276		PBS	
		Alive/treated	Survival %	Alive/treated	Survival %
JM9130013	VIII	nd	nd	nd	nd
CJB111	V	47/49	96	12/46	26
COH1	III	47/50	94	12/50	24
M781	III	33/50	66	6/50	12
2603	V	28/30	93	8/48	17
18RS21	II	31/78	40	10/46	22
DK21	II	37/68	54	15/60	25
H36B	Ib	8/38	21	5/60	8
7357b –	Ib	29/50	58	5/50	10
A909	Ia	18/49	37	6/49	12

Accordingly, the invention therefore includes compositions comprising combinations of two or more GBS antigens, wherein the combination includes GBS 80 or a fragment thereof or a 10 polypeptide sequence having sequence identity thereto.

In one embodiment, the combination may consist of two to thirteen GBS antigens, including GBS 80. As an example, the combination may contain GBS 80 and other GBS antigens selected from the group consisting of GBS 80, GBS 91, GBS 104, GBS 184, GBS 276, GBS 305, GBS 322, GBS 330, GBS 338, GBS 361, GBS 404, GBS 690, and GBS 691. Preferably, the combination includes

GBS 80 in combination with one or more of GBS 104 and GBS 322. For example, the combination may include GBS 80, GBS 104, GBS 322 and GBS 67.

Instead of the full length antigen, the combination may comprise an immunogenic fragment of the selected GBS antigen and/or a polypeptide sequence having sequence identity to the selected antigen.

Preferably, the combination of GBS antigens consists of three, four, five, six, seven, eight, nine, or ten GBS antigens. Still more preferably, the combination of GBS antigens consists of three, four, or five GBS antigens.

Details of examples of GBS antigens for use in combination with GBS 80 are set forth below.

10 **GBS 91**

GBS 91 refers to a GBS C3 binding polypeptide. Nucleotide and amino acid sequences of GBS 91 sequenced from serotype V isolated strain 2603 V/R are set forth in Ref. 2 as SEQ ID 8937 and SEQ ID 8938. These sequences are set forth below as SEQ ID NOS 10 and 11:

15 **SEQ ID NO. 10**

ATGAAAAAAGGACAAGTAAATGATACTAAGCAATCTTACTCTACGTAAATATAAATTGGTTAGC
 ATCAGTAATTAGGGTCATTCTATAATGGTCACAAGTCCTGTTTGGATCAAACATCGGTTCAAGT
 AAGTTAATAATCAGACAGGCAGTAGTGTTGATGCTAATAATTCTCCAATGAGACAAGTGCCTCAAGT
 GTGATTACTTCCAATAATGATAGTGTCAAGCGCTGTATAAAGTTGTAATAAGTCAAAATACGGCAAC
 20 AAAGGACATTACTACTCCTTAGTAGAGACAAAGCCAATGGTGGAAAAAACATTACCTGAACAGGGA
 ATTATGTTATAGCAAAGAAACCGAGGTGAAAAATACACCTTCAAAATCAGCCCCAGTAGCTTCTAT
 GCAAAGAAAGGTGATAAAAGTTCTATGACCAAGTATTAATAAGATAATGTGAAATGGATTTCATA
 TAAGTCTTTGTGGCTACGTCGATACGCAGCTATTGAGTCAGTACAGATCCATCAGGAGGTTCAGAGA
 25 CTAAAGCACCTACTCCTGTAACAAATTAGGAAGCAATAATCAAGAGAAAATAGCAACGCAAGGAAAT
 TATACATTTCACATAAAAGTAGAAGTAAAAATGAAGCTAAGGTAGCGAGTCACACTCAATTACATT
 GGACAAAGGAGACAGAATTTTACGACCAAATACTAACTATTGAAGGAAATCAGTGGTTATCTTATA
 AATCATTCAATGGTGTCTCGTTGCTAGTAAAGCATCTCAGTAGAGAAAAACTGAAGAT
 30 AAAGAAAAAGTGTCTCCTCAACCACACAAGCCGTATTACTAAAGACTGGTAGACTGACTATTCTAACGA
 AACAACTACAGGTTTGATATTAAATTACGAATATTAAAGATGATAACGGTATCGCTGCTGTTAAGG
 TACCGGTTGGACTGAACAAGGAGGGCAAGATGATATTAAATGGTATACAGCTGTAACACTGGGGAT
 GGCAACTACAAAGTAGCTGTATCATTGCTGACCATAAGAATGAGAAGGGTCTTATAATATTCAATT
 ATACTACCAAGAAGCTAGTGGGACACTGTAGGTGTAACAGGAACCTAAAGTGACAGTAGCTGGAAC
 ATTCTCTCAAGAACCTATTGAAATGGTTAGCAAAGACTGGTGTATAATATTATCGGAAGTACT
 35 GAAGTAAAAATGAAGCTAAATATCAAGTCAGACCAATTACTTAGAAAAAGGTGACAAAATAAA
 TTATGATCAAGTATTGACAGCAGATGGTACCAAGTGGATTCTACAAATCTATAGTGGTGTTCGTC
 GCTATATTCTGTAAAAAGCTAACTACAAGTAGTGAAAGCGAAAGATGAGGCGACTAAACCGACT
 AGTTATCCCAACTTACCTAAACAGGTACCTATACATTACTAAACTGTAGATGTGAAAGTCAACC
 TAAAGTATCAAGTCCAGTGGATTAAATTCAAAAGGTGAAAAAATACATTATGATCAAGTGTAG
 40 TAGTAGATGGTCATCAGTGGATTTCATACAAGAGTTATTCCGGTATTGTCGCTATATTGAAATT

SEQ ID NO. 11

MKKQVNDTKQSYSLRKYKFGLASVILGSFIMVTPVFADQTTSVQVNNQTGTSVDANNSSNETSASS
VITSNNDSVQASDKVVNSQNTATKDITPLVETKPMVEKTLPEQGNVYSKETEVKNTPSKSAPVAFY
AKKGDKVFYDQVFNKDNVKWISYKSFCGVRRYAAIESLDPSGGSETKAPTPVTNSGSNNQEKIATQGN
YTFSHKVEVKNEAKVASPTQFTLDKGDRIFYDQILTIEGNQWL SYKS FNGVRRFVLLGKASSVEKTED
KEKVSPQPQARITKTGRLTISNETTGFDILITNIKDDNGIAAVKVPVWTEQGGQDDIKWYTAVTTGD

GNVKVAVS FADHKNEKGLYNIHLYYQEASGTLVGVGTGVTVAGTNSSQEPIENGLAKTGVYNIIGST
EVKNEAKI SQTQFTLEKGDKINYDQVLTADGYQWISYKSYSGVRRYIPVKLTTSEKAKDEATKPT
SYPNL PKTGTYTFTKTVDVKSQPKVSSPVEFN FQKGEKIHYDQVLVVDGHQWISYKSYSGIRRYIEI

5 GBS 91 contains an N-terminal leader or signal sequence region which is indicated by the underlined sequence at the beginning of SEQ ID NO: 11 above. In one embodiment, one or more amino acids from this leader or signal sequence region of GBS 91 are removed. An example of such a GBS 91 fragment is set forth below as SEQ ID NO: 12.

10 **SEQ ID NO: 12**

DQTTSVQVNNQTGTSVDANNSSNETSASSVITSNNDSVQASDKVVNSQNTATKDI TTPLVETKPMVEK
TLPEQGNVYVSKETEVKNTPSKSAPVAFYAKKGDKVFYDQVFNKDNVKWISYKSFCGVRRYAAIESLD
PSGGSETKAPTPTVNSGSNNQEKIATQGNYTFSHKVEVKNEAKVASPTQFTLDKGDRIFYDQILTIEG
NQWLSYKSFNGVRRFVLLGKASSVEKTEDKEKVS PQPQARITKTGRLTISNETTGF DILITNIKDDN
15 GIAAVKVPVWTEQGGQDDIKWYTAVTTGDGNYKVAVS FADHKNEKGLYNIHLYYQEASGTLVGVGTGK
VTVAGTNSSQEPIENGLAKTGVYNIIGSTEVKNEAKI SQTQFTLEKGDKINYDQVLTADGYQWISYK
SYSGVRRYIPVKLTTSEKAKDEATKPT SYPNL PKTGTYTFTKTVDVKSQPKVSSPVEFN FQKGEKI
HYDQVLVVDGHQWISYKSYSGIRRYIEI

20 GBS 91 contains a C-terminal transmembrane region which may be located within the underlined region near the end of SEQ ID NO: 11 above. In one embodiment, one or more amino acids from the transmembrane and cytoplasmic regions are removed. An example of such a GBS 91 fragment is set forth below as SEQ ID NO: 13.

25 **SEQ ID NO: 13**

MKKGQVNDTKQSYSLRKYKFGLASVILGSFIMVTS PVFADQTTSVQVNNQTGTSVDANNSSNETSASS
VITSNNDSVQASDKVVNSQNTATKDI TTPLVETKPMVEK TLPEQGNVYVSKETEVKNTPSKSAPVAFY
AKKGDKVFYDQVFNKDNVKWISYKSFCGVRRYAAIESLDPSGGSETKAPTPTVNSGSNNQEKIATQGN
30 YTFSHKVEVKNEAKVASPTQFTLDKGDRIFYDQILTIEGNQWLSYKSFNGVRRFVLLGKASSVEKTED
KEKVSPQPQARITKTGRLTISNETTGF DILITNIKDDNGIAAVKVPVWTEQGGQDDIKWYTAVTTGD
GNYKVAVS FADHKNEKGLYNIHLYYQEASGTLVGVGTGVTVAGTNSSQEPIENGLAKTGVYNIIGST
EVKNEAKI SQTQFTLEKGDKINYDQVLTADGYQWISYKSYSGVRRYIPVKLTTSEKAKDEATKPT
SYPNL PKTG

35 GBS 91 contains an amino acid motif indicative of a cell wall anchor: **SEQ ID NO: 14**
LTKTG (shown in italics in SEQ ID NO: 11 above). In one embodiment, both the transmembrane domain and the cell wall anchor motif are removed from GBS 91. An example of such a GBS 91 fragment is set forth below as SEQ ID NO: 15.

40 **SEQ ID NO: 15**

MKKGQVNDTKQSYSLRKYKFGLASVILGSFIMVTS PVFADQTTSVQVNNQTGTSVDANNSSNETSASS
VITSNNDSVQASDKVVNSQNTATKDI TTPLVETKPMVEK TLPEQGNVYVSKETEVKNTPSKSAPVAFY
AKKGDKVFYDQVFNKDNVKWISYKSFCGVRRYAAIESLDPSGGSETKAPTPTVNSGSNNQEKIATQGN
45 YTFSHKVEVKNEAKVASPTQFTLDKGDRIFYDQILTIEGNQWLSYKSFNGVRRFVLLGKASSVEKTED
KEKVSPQPQARITKTGRLTISNETTGF DILITNIKDDNGIAAVKVPVWTEQGGQDDIKWYTAVTTGD
GNYKVAVS FADHKNEKGLYNIHLYYQEASGTLVGVGTGVTVAGTNSSQEPIENGLAKTGVYNIIGST

EVKNEAKISSQTQFTLEKGDKINYDQVLTADGYQWISYKSYSVRRYIPVKLTTSEKAKDEATKPT
SYPN

5 In one embodiment, one or more amino acids from the leader or signal sequence region and
one or more amino acids from the transmembrane and cytoplasmic regions are removed from the GBS
91 sequence. An example of such a GBS 91 fragment is set forth below as SEQ ID NO: 16.

SEQ ID NO: 16

10 DQTTSVQVNNQTGTSVDANNSSNETSASSVITSNNDSVQASDKVVNSQNTATKDITTPVETKPMVEK
TLPEQGNYVYSKETEVKNTPSKSAPVAFYAKKGDKVFYDQVFNKNVWISYKFCGVRRYAAIESLD
PSGGSETKAPTPVTNSGSNNQEKIATQGNYTFSHKVEVKNEAKVASPTQFTLDKGDRIFYDQILTIEG
NQWLSYKSFNGVRRFVLLGKASSVEKTEDKEKVSPQPQARITKTGRLTISNETTGFDILITNIKDDN
GIAAVKVPVWTEQGGQDDIKWYTAVTTGDGNYKVAVSFADHKNEKGLYNIHLYYQEASGTLVGVGTGK
15 VTVAGTNSSQEPIENGLAKTGVYNIIGSTEVKNEAKISSQTQFTLEKGDKINYDQVLTADGYQWISYK
SYSVRRYIPVKLTTSEKAKDEATKPTSYPNLPKTG

In another embodiment, the leader or signal sequence region, the transmembrane and
cytoplasmic regions, and the cell wall anchor motif are all removed from the GBS 91 sequence. An
example of such a GBS 91 fragment is set forth below as SEQ ID NO: 17.

20

SEQ ID NO: 17

25 DQTTSVQVNNQTGTSVDANNSSNETSASSVITSNNDSVQASDKVVNSQNTATKDITTPVETKPMVEK
TLPEQGNYVYSKETEVKNTPSKSAPVAFYAKKGDKVFYDQVFNKNVWISYKFCGVRRYAAIESLD
PSGGSETKAPTPVTNSGSNNQEKIATQGNYTFSHKVEVKNEAKVASPTQFTLDKGDRIFYDQILTIEG
NQWLSYKSFNGVRRFVLLGKASSVEKTEDKEKVSPQPQARITKTGRLTISNETTGFDILITNIKDDN
GIAAVKVPVWTEQGGQDDIKWYTAVTTGDGNYKVAVSFADHKNEKGLYNIHLYYQEASGTLVGVGTGK
VTVAGTNSSQEPIENGLAKTGVYNIIGSTEVKNEAKISSQTQFTLEKGDKINYDQVLTADGYQWISYK
30 SYSVRRYIPVKLTTSEKAKDEATKPTSYPN

30

Further information regarding GBS 91 can be found in WO 01/25440 (C3 binding
polypeptide), WO 01/32882 (ID-65), WO 02/31156 (BVH) and Reinscheid et al., *Microbiology*
(2002) 148: 3245-3254 (*bsp* gene), each of which are incorporated herein by reference in their
entirety.

35

GBS 104

GBS 104 refers to a putative cell wall surface anchor family protein. It has been referred to as
emaA protein. Nucleotide and amino acid sequences of GBS 104 sequenced from serotype V isolated
strain 2603 V/R are set forth in Ref. 2 as SEQ ID 8777 and SEQ ID 8778. These sequences are set
forth below as SEQ ID NOS 18 and 19:

SEQ ID NO. 18

ATGAAAAAGAGACAAAAAATATGGAGAGGGTTATCAGTTACTTACTAATCCTGCCCCAATTCCATT
 5 TGGTATATTGGTACAAGGTGAAACCCAAGATAACCAATCAAGCACTGGAAAAGTAATTGTTAAAAAAA
 CGGGAGACAAATGCTACACCATTAGGCAAAGCGACTTGTGTTAAAAAATGACAATGATAAGTCAGAA
 ACAAGTCACGAAACGGTAGAGGGTCTGGAGAAGCAACCTTGAAAACATAAAACCTGGAGACTACAC
 ATTAAGAGAAGAAACAGCACCAATTGGTTATAAAAACACTGATAAAACCTGGAAAGTTAAAGTTGCAG
 10 ATAACGGAGCAACAATAATCGAGGGTATGGATGCAGATAAAGCAGAGAAACGAAAAGAAGTTGAAT
 GCCCAATATCCAAAATCAGCTATTATGAGGATACAAAAGAAAATTACCCATTAGTTAATGTAGAGGG
 15 TTCCAAAGTTGGTGAACAATACAACAGCATTGAATCCAATAATGGAAAAGATGGTCGAAGAGAGATTG
 CTGAAGGTTGGTTATCAAAAAAAATTACAGGGGTCATGATCTCGATAAGAATAAATATAAAATTGAA
 TTAACTGTTGAGGGTAAACCACTGTTGAAACGAAAGAACTTAATCAACCACTAGATGTCGTTGCT
 ATTAGATAATTCAAATAGTATGAATAATGAAAGAGCCAATAATTCTCAAAGAGCATTAAAGCTGGGG
 20 AAGCAGTTGAAAAGCTGATTGATAAAATTACATCAAATAAAGACAATAGAGTAGCTCTGTGACATAT
 GCCTCAACCATTGATGGTACTGAAGCGACCGTATCAAAGGGAGTTGCCGATCAAATGGTAAAGC
 GCTGAATGATAGTGTATCATGGGATTATCATAAAACTACTTTACAGCAACTACACATAATTACAGTT
 ATTAAATTAAACAAATGATGCTAACGAGTTAATATTCTAAAGTCAGAAATTCCAAGGAAGCAGGGAG
 25 CATATAATGGGGATCGCACGCTCTATCAATTGGTGCACATTACTCAAAAGCTCTAATGAAAGC
 AAATGAAATTAGAGACACAAAGTTCTAATGCTAGAAAAAAACTTATTTACGTAACTGATGGT
 30 TCCCTACGATGTCTTATGCCATAAATTAACTCCTTATATCAACATCTTACCAAAACCAGTTAAT
 TCTTTTTAAATAAAATACCAGATAGAAGTGGTATTCTCAAAGAGGATTTTATAATCAATGGTGTG
 TTATCAAATAGTAAAGGAGATGGAGAGGTTAAACTGTTTCGGATAGAAAAGTTCTGTACTG
 GAGGAACGACACAACGAGCTTATCGAGTACCGAAAATCAACTCTCTGTAATGAGTAATGAGGGATAT
 GCAATTAAATAGTGGATATATTATCTCTATTGGAGAGATTACAACGGTCTATCCATTGATCCTAA
 35 GACAAAGAAAGTTCTGCAACGAAACAAATCAAACACTCATGGTGGAGCAACACATTATACTTTAATG
 GAAATATAAGACCTAAAGTTATGACATTTTACTGTTGGGATTGGTGTAAACGGAGATCCTGGTGCA
 ACTCCTCTGAAGCTGAGAAATTATGCAATCAATATCAAGTAAACAGAAAATTATACTAATGTTGA
 TGATACAAATAAAATTATGATGAGCTAAATAACTTTAAACAAATTGTTGAGGAAAACATTCTA
 TTGTTGATGAAATGTGACTGATCCTATGGGAGAGATGATTGAATTCCAATTAAAAATGGTCAAAGT
 40 TTTACACATGATGATTACGTTGGTGGAAATGATGGCAGTCATTAAAAATGGTGTGGCTCTGG
 TGGACCAACAGTGTGGGGATTAAAGATGTTACAGTGACTTATGATAAGACATCTCAAACCA
 TCAAAATCAATCATTGAACCTAGGAAGTGGACAAAAGTAGTTCTACCTATGATGTACGTTAAAA
 GATAACTATATAAGTAAACAAATTTCACAATACAATAATCGTACAACGCTAAGTCCGAAGAGTAAAA
 AGAACCAAATACTATCGTGTGATTCCCAATTCCAAAATTGATGTTGAGTTCCGGTACTAA
 45 CCATCAGTAATCAGAAGAAAATGGGTGAGGTGAATTATTAAAGTTAATAAAGACAAACATTAGAA
 TCGCTTTGGGAGCTAAGTTCAACTTCAGATAGAAAAAGATTCTGGGTATAAGCAATTGTTCC
 AGAGGGAAAGTGTACAACAAAGAATGATGGTAAATTATTTAAAGCACTTCAGATGGTAAC
 ATAAATTATGAAATTCAAGTCCAGATGGCTATATAGAGGTTAAAACGAAACCTGTTGACATT
 ACAATTCAAATGGAGAAGTTACGAACCTGAAAGCAGATCCAATGCTAATAAAATCAAATCGGGTA
 50 TCTTGAAGGAAATGGTAAACATCTTATTACCAACACTCCCAACGCCACCAGGTGTTTCTAAAA
 CAGGGGAAATTGGTACAATTGCTATATATTAGTTGGTTACTTTATGATACTTACCAATTGTTCT
 TTCCGTGTAACAAATTG

SEQ ID NO. 19

45 MKKRQKIWRGLSVTLLILSQIPFGILVQGETQDTNQALGKVIVKKTGDNATPLGKATFVLKNDNDKSE
 TSHETVEGSGEATFENIKPGDYTLREETAPIGYKKTDKTWKVKVADNGATIIEGMDADKAERKEVLN
 AQYPKSAIYEDTKENYPLVNVEGSKVGEQYKALNPINGKDRREIAEGWLSKKITGVNDLKDKNKYKIE
 LTVEGKTTVETKELNQPLDVVVLLDNSNSMNNEARRANSQRALKAGEAVEKLIKITSNKDNRVALVTY
 ASTIFDGTEATVSKGVADQNGKALNDSVSWDYHKTTFTATTHNYSYLNLTNDANEVNILKSRIPKEAE
 50 HINGDRTLYQFGATFTQKALMKANEILETQSSNARKKLIFHVTDGVPMSYAINFNPYISTSYQNFN
 SFLNKIPDRSGILQEDFIINGDDYQIVKGDGESFKLFSDRKVPVTGGTTQAAYRVPQNQLSVMNSNEG
 AINSGYIYLWRYDYNWVYPFDPKTKVVSATKQIKTHGEPTTLYFNGNIRPKGYDIFTVGIGVNGDPGA
 TPLEAEKFMQSISSKTENYTNVDDTNKIYDELNKYFKTIVEEKHSIVDGNVTDPGMEMIEFQLKNGQS

5

FTHDDYVLVGNDGSQLKNGVALGGPNSDGGILKDVTVTYDKTSQTIKINHNLGSGQKVVLTYDVRLK
 DNYISNKFYNTNNRTTLSPKSEKEPNTIRDFPIPKIRDVREFPVLTISNQKKMGEVEFIKVNKDKHSE
 SLLGAKFQLQIEKDFSGYKQFVPEGSDVTTKNDGKIYFKALQDGNYKLYEISSPDGYIEVTKPVVTF
 TIQNGEVTNLKADPNANKNQIGYLENGKHLITNTPKRPPGVFPKTGGIGTIVYILVGSTFMILTICS
FRRKQL

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GBS 104 contains an N-terminal leader or signal sequence region which is indicated by the underlined sequence at the beginning of SEQ ID NO 19 above. In one embodiment, one or more amino acid sequences from the leader or signal sequence region of GBS 104 are removed. An example of such a GBS 104 fragment is set forth below as SEQ ID NO 20.

SEQ ID NO 20

GETQDTNQALGKIVVKKTGDNATPLGKATFVLKNDNDKSETSHETVEGSGEATFENIKPGDYTLREET
 APIGYKKTDKTWKVVKVADNGATIIEGMDADKAEKRKEVLNAQYPKSAIYEDTKENYPLVNVEGSKVGE
 15 QYKALNPINGKDGRREIAEGWLSKKITGVNDLDKNKYKIELTVEGKTTVETKELNQPLDVVVLDDNSN
 SMNNERANNSQRALKAGEAVEKLIKDITSNKDNRVALVTYASTIFDGTEATVSKGVADQNGKALND
 SWDYHKTTFTATTHNSYLNLTNDANEVNILKSRIPKAEAHINGDRTLYQFGATFTQKALMKANEILE
 20 TQSSNARKKLIFHVTDGVPVMSYAINFNPYISTSYQNQFNSFLNKIPDRSGILQEDFIINGDDYQIVK
 GDGESFKLFSDRKVPVTTQAAAYRVPQNQLSVMSEN
 25 ATKQIKTHGEPTTLYFNGNIRPKGYDIFTVGIGVNGDPGATPLEAEKFMQSISSSKTE
 YDELNKYFKTIVEEKHSIVDGNVTDPGEMIEFQLKNGQSFTHDDYVLVGNDSQLKNGVALGGPNSD
 GGLKDVTVTYDKTSQTIKINHNLGSGQKVVLTYDVRKDNYISNKFYNTNNRTTLSPKSEKEPNTI
 RDPIPKIRDVREFPVLTISNQKKMGEVEFIKVNKDKHSESSLLGAKFQLQIEKDFSGYKQFVPEGSDV
 TTKNDGKIYFKALQDGNYKLYEISSPDGYIEVTKPVVTFTIQNGEVTNLKADPNANKNQIGYLENG
 25 KHLITNTPKRPPGVFPKTGGIGTIVYILVGSTFMILTICSFRRKQL

30

GBS 104 contains a C-terminal transmembrane and/or cytoplasmic region which is indicated by the underlined region near the end of SEQ ID NO 19 above. In one embodiment, one or more amino acids from the transmembrane or cytoplasmic regions are removed. An example of such a GBS 104 fragment is set forth below as SEQ ID NO 21.

SEQ ID NO: 21

MKMRQKIWRGLSVTLLILSQIPFGILVQGETQDTNQALGKIVVKKTGDNATPLGKATFVLKNDNDKSE
 35 TSHETVEGSGEATFENIKPGDYTLREETAPIGYKKTDKTWKVVKVADNGATIIEGMDADKAEKRKEVLN
 AQYPKSAIYEDTKENYPLVNVEGSKVGEQYKALNPINGKDGRREIAEGWLSKKITGVNDLDKNKYKIE
 LTVEGKTTVETKELNQPLDVVVLDDNSNSMNNEARRANSQRALKAGEAVEKLIKDITSNKDNRVALVTY
 ASTIFDGTEATVSKGVADQNGKALNDSVWDYHKTTFTATTHNSYLNLTNDANEVNILKSRIPKAE
 HINGDRTLYQFGATFTQKALMKANEILETQSSNARKKLIFHVTDGVPVMSYAINFNPYISTSYQNQFN
 40 SFLNKIPDRSGILQEDFIINGDDYQIVKGDGESFKLFSDRKVPVTTQAAAYRVPQNQLSVMSEN
 AINSGYIYLYWRDYNWVYPFDPKVVSATKQIKTHGEPTTLYFNGNIRPKGYDIFTVGIGVNGDPGA
 TPLEAEKFMQSISSSKTE
 45 YNTNVDDTNKIAEDLNKYFKTIVEEKHSIVDGNVTDPGEMIEFQLKNGQSFTHDDYVLVGNDSQLKNGVALGGPNSDGGILKDVTVTYDKTSQTIKINHNLGSGQKVVLTYDVRKDNYISNKFYNTNNRTTLSPKSEKEPNTI RDPIPKIRDVREFPVLTISNQKKMGEVEFIKVNKDKHSESSLLGAKFQLQIEKDFSGYKQFVPEGSDV TTKNDGKIYFKALQDGNYKLYEISSPDGYIEVTKPVVTFTIQNGEVTNLKADPNANKNQIGYLENG KHLITNTPKRPPGVFPKTGGIGTIVYILVGSTFMILTICSFRRKQL

In one embodiment, one or more amino acids from the leader or signal sequence region and one or more amino acids from the transmembrane or cytoplasmic regions are removed. An example of such a GBS 104 fragment is set forth below as SEQ ID NO 22.

5 **SEQ ID NO: 22**

GETQDTNQALGKVIVKKTGDNATPLGKATFVLKNDNDKSETSHETVEGSGEATFENIKPGDYTLREET
 APIGYKKTDKTWKVKVADNGATIIEGMDADKAERKEVLNAQYPKSAIYEDTKENYPLVNVEGSKVGE
 QYKALNPINGKDRREIAEGWLSKKITGVNDLKDKNKYKIELTVEGKTTVETKELNQPLDVVLLDNSN
 10 SMNNERANNSQRALKAGEAVEKLIDKITSNKDNRVALVTYASTIFDGTEATVSKGVADQNGKALNDSV
 SWDYHKTTFTATTHNYSYLNLTNDANEVNILKSRIPKAEAHINGDRTLYQFGATFTQKALMKANEILE
 TQSSNARKKLIFHVTDGVPTMSYAINFNPYISTSYQNQFNSFLNKIPDRSGILQEDFIINGDDYQIVK
 GDGESFKLFSDRKVPVTGGTTQAAYRVPQNQLSVMNSNEYAINGSYIYLYWRDYNWVYPFDPKTKV
 15 ATKQIKTHGEPTTLYFNGNIRPKGYDIFTVGIGVNGDPGATPLEAEKFMQSISSTENYTNVDDTNKI
 YDELNKYFKTIVEEKHSIVDGNVTDPGMEMIEFQLKNGQSFTHDDYVLVGNNDGSQLKNGVALGGPN
 20 SDGGILKDVTVTYDCTSQTIKINHLNGSQKVVLTYDVRLLDNYISNKFYNTNNRTTLSPKSEKEPNTI
 RDFPIPKIRDVREFPVLTISNQKKMGEVEFIKVNKDKHSESLLGAKFQLQIEKDFSGYKQFVPEGSDV
 TTKNDGKIYFKALQDGNYKLYEISSPDGYIEVKTKPVVTFTIQNGEVTNLKADPNANKNQIGYLEGNG
 KHLITNT

20 In other embodiments, additional fragments of GBS 104 are provided including an 830 amino acid fragment of GBS 104 of amino acids 28-858, a 359 amino acid fragment of GBS 104 of amino acids 28-387, a 581 amino acid fragment of GBS 104 of amino acids 28-609, or a 740 amino acid fragment of GBS 104 of amino acids 28-768.

25 **GBS 184**

GBS 184 refers to a putative lipoprotein. Nucleotide and amino acid sequences of GBS 184 sequenced from serotype V isolated strain 2603 V/R are set forth in Ref. 2 as SEQ ID 1977 and SEQ ID 1978. These sequences are also set forth below as SEQ ID NOS 23 and 24.

30 **SEQ ID NO: 23**

ATGAAAAAAACAAAAACTATTACTGCTTATTGGAGGCTTATTAATAATGATAATGATGACAGCATGTA
 GGATTCAAAAATCCCAGAAAACCGCACAAAGGAAGAGTACCAAGCTGAACAAAATTAAACCGTT
 35 TTGAGTTTTAGCACAAAAGATAAAGATTGAGCAAAATACAAAATACTTACTATTAGTATCGGAT
 TCAGGTGATGCATTAGATTAGAATATTCTATAGTATTCAAGATTAAAAAAAATAAGGATTAGG
 GAAGTTGAAACAAGAAAAGTCAAATAGAAAAGCCGGTGGCTATAATGAGTTAGAAAATAAGAGG
 TCCCATTGAAATATTAAAAATAATATAGTTATCCAAAAGGAAAACCGAATATTACATTGATGAC
 40 TTTATTATCGGAGCAATGGATACTAAAGAATTAAAAGAATTAAAAAAATTAAAAGTAAAAGTTATT
 ATTAACATCCGGAACTGAGTTGAAAGATATAACATATGAATTGCCGACACAGTCGAAGCTTATTA
 AAAAA

SEQ ID NO: 24

MKKQKLLLLIGGLLIMIMMTACKDSKIPENRTKEEYQAEQNFKPFFFLAQKDKDLSKIQKYLLLVSD
 SGDALDLEYFYSIQDLKKNLDGFETRKSQIEKPGGYNELENKEVPFEYFKNNIVYPKGKPNTFDD
 45 FIIGAMDTKELKELKKLKVKSYLLKHPETELKDTYELPTQSKLIKK

GBS 184 contains a N-terminal leader or signal sequence region which is indicated by the underlined sequence at the beginning of SEQ ID NO 24, above. In one embodiment, one or more amino acids from the leader or signal sequence are removed from GBS 184. An example of such a GBS 184 fragment is set forth below as SEQ ID NO: 25.

5

SEQ ID NO: 25

KDSKIPENRTKEEYQAEQNFKPFFEFLAQKDKDLSKIQKYLLLVSDSGDALDLEYFYSIQDLKKNKL
GKFETRKSQIEKPGGYNELENKEVPFYEYFKNNIVYPKGKPNITFDDFIIGAMDTKELKELKKLVKSY
LLKHPETELKDITYELPTQSKLIKK

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GBS 276

GBS 276 refers to a C5a peptidase. Nucleotide and amino acid sequences of GBS 276 sequenced from serotype V isolated strain 2603 V/R are set forth in Ref. 2 as SEQ ID 8941 and SEQ ID 8942. These sequences are set forth below as SEQ ID NOS 26 and 27:

15

SEQ ID NO. 26

TTGC GTAAAAAAACAAAAACTACCATTGATAAAACTTGCCTATTGCCATTGCGCTTATATCTACGAGCATCTTGCT
CAATGCACAAATCAGACATTAAAGCAAATACTGTGACAGAAAGACACTCCTGCTACCGAACAGCCGTAG
AACC CCCACAACCAATAGCAGTTCTGAGGAATCAGATCATCAAAGGAACTAAAACCTCACAAACT
20 CCTAGTGATGTAGGAGAAACAGTAGCAGATGACGCTAATGATCTAGCCCCCTCAAGCTCCTGCTAAAAC
TGCTGATAACACCAGCAACCTCAAAGCGACTATTAGGGATTGAACGACCCCTCTCATGTCAAAACCC
TGCAAGGAAAAGCAGGCAAGGGAGCTGGGACCGTTGTCAGTGATGCTGGTTTGATAAAAAT
CATGAAGCGTGGCGCTTAACAGACAAAACTAAAGCACGTTACCAATCAAAGAAAATCTGAAAAAGC
25 TAAAAAAGAGCACGGTATTACCTATGGCGAGTGGGTCATGATAAGTTGCTTATTACACGACTATA
GTAAAGATGGTAAAACGCTGTTGATCAAGAACACGGCACACACGTGTCAGGGATCTTGTCAAGGAAT
GCTCCATCTGAAATGAAAGAACCTTACCGCCTAGAAGGTGCGATGCCTGAGGCTCAATTGCTTTGAT
GCCTGTCGAAATTGTAATGGACTAGCAGACTATGCTCGTAACTACGCTCAAGCTATCAGAGATGCTG
TCAACTGGGAGCTAAGGTGATTAATATGAGCTTGGTAATGCTGCACTAGCTTACGCCAACCTTCCA
30 GACGAAACCAAAAAGCCTTGTACTATGCCAAATCAAAGGTGTTAGCATTGACCTCAGCTGGTAA
TGATAGTAGCTTGGGGCAAGCCCCGCTACCTCTAGCAGATCATCCTGATTATGGGGTGGTTGGGA
CACCTGCAGCGGCAGATTCAACATTGACAGTTGCTTCTACAGCCCAGATAAACAGCTCACTGAAACT
GCTACGGTAAAACAGACGATCATCAAGATAAAGAAATGCCTGTTATTCAACAAACCGTTTGAGCC
AAACAAGGCTTACGACTATGCTTATGCTAATCGTGGTACGAAAGAGGATGATTAAAGGATGTCGAAG
35 GTAAGATTGCCCTTATTGAACGTGGCGATATTGATTCAAAGATAAGATTGCAAACGCTAAAAAGCT
GGTGTGAGGGTCTGATCTAGACAACTCAAGACAAGGGCTCCGATTGAATTGCCAAATGTTGA
CCAGATGCCCTGCCCTTATCAGTCGAAGAGACGGCTCTTATTAAAAGACAATCCCCAAAAACCA
TTACCTCAATGCGACACCTAAGGTATTGCCAACAGCAAGTGGCACCAAACATAAGCCGTTCTCAAGC
40 TGGGGTCTGACAGCTGACGGCAATTAAACCGGATATTGCAAGCACCCGGCAAGATAATTGTCATC
AGTGGCTAACACAAGTATGCAAACATTCTGGAACTAGTATGTCGACCAATTGGTAGCGGGTATCA
TGGGACTGTTGCAAAGCAATATGAGACACAGTATCCTGATATGACACCACAGAGCGTCTTGATT
GCTAAGAAAGTATTGATGAGCTAGCAACTGCCCTATATGATGAAGATGAAAAGCTTATTTCTCC
TCGCCAACAGGGAGCAGGAGCAGTCGATGCTAAAAAGCTCAGCACGAAACGATGTATGAAACAGATA
AGGACAATACCTCAAGCAAGGTTCACCTGAACAAATGTTGATAAATTGAGTAACAGTAACAGTT
45 CACAAACAAATCTGATAAACCTCAAGAGTTGATTACCAAGTAACGTTCAAACAGATAAAGTAGATGG
AAAACACTTGCCTGGCTCTAAAGCATTGATGAGACATCATGGCAAAAATCACAATTCCAGCCA
ATAGCAGCAAACAAGTCACCCTCCAATCGATGCTAGTCGATTAGCAAGGACTTGCTGCCAAATG
AAAAATGGCTATTCTTAGAAGGTTGGTCAAAACAAGATCCTACAAAAGAAGAGCTTATGAG
CATTCATATATTGGTTCCGAGGTGATTGGCAATCTGTCAGCCTAGAAAACCAATCTATGATA

GCAAAGACGGTAGCAGCTACTATCATGAAGCAAATAGTGATGCCAAGACCAATTAGATGGTATGGA
 TTACAGTTTACGCTCTGAAAATAACTTACAGCACTTACACAGAGTCTAACCCATGGACGATTAT
 TAAAGCTGTCAAAGAAGGGGTGAAAACATAGAGGATATCGAATCTCAGAGATCACAGAAACCATT
 TTGCAGGTACTTTGCAAACAAAGACGATGATGCCACTACTATATCCACCGTCACGCTAATGGAAA
 5 CCATATGCTCGATCTCTCAAATGGGACGGTAACAGAGATTATGCTCAATTCAAGGTACTTCTT
 GCGTAA TGCTAAAACCTTGTGGCTGAAGTCTGGACAAAGAAGGAATGTTGTTGGACAAGTGAGG
 TAACCGAGCAAGTTAAAACATACAACAAATGACTTGGCAAGCACACTGGTTCAACCGTTGAA
 AAAACCGCTGGGACGGTAAAGATAAAAGACGGCAAAGTTGCTAACGGAACCTACACCTATCGTGT
 TCGCTACACGCCGATTAGCTAGGTGCAAAGAACACACACTGATTTGATGATTGAGACAATA
 10 CGACACCTGAAGTCGCAACATCGGCAACATTCTAACAGAACAGATAGTCGTTGACACTGACATCTAAA
 CCAAAACCAACGGTTACCGTGAGCGTATTGCTTACACTTATGGATGAGGATCTGCCAAC
 AACAGAGTATATTCTCAAATGAAGATGGTACCTTACTCTTCTGAAGAGGCTGAAACAATGAAAG
 GCGCTACTGTTCCATTGAAAATGTCAGACTTACTTATGTTGAAGATATGGCTGGTAACATCACT
 TATACACCAGTGACTAAGCTATTGGAGGGCCACTCTAATAAGCCAGAACAGACGGTCAGATCAAGC
 15 ACCAGACAAAGAAACCAAGCTAAACCAAGAACAGCGGTCAGGTCAAACACCAGATAAAAAAAAG
 AAACTAAACCAGAAAAAGATAGTCAGGTCAAACACCAGGTAACCTCTCAAAAGGTCAATCTCT
 CGTACTCTAGAGAAACGATCTCTAACCGTGCTTAGCTACAAAAGCATCAACAAGAGATCAGTTAC
 AACGACTAATGACAAGGATAAAATGTTACATCTCTTAAGTTAGTTATGACCACCTTCTTGG
 20 GA

SEQ ID NO. 27

MRKKQKL~~PF~~DKLAI~~AL~~LISTSILLNAQSDIKANTVTE~~D~~PATEQAVEPPQPIAVSEESRSSKETKTSQT
 PSDVGETVADDANDIAPQAPAKTADTPATSKATIRDLNDPSHVKT~~I~~QEKAGKGAGTVVAVIDAGFDKN
 HEAWRLTD~~K~~T~~K~~ARYQS~~K~~ENLEKAKKEHGITYGEW~~V~~NDKVAYYHDYSKDGN~~A~~VDQEHGTHVSGILSGN
 25 APSEMKE~~P~~YRLEGAMPEAQLL~~M~~RVEIVNGLADYARNYQAIRDAVN~~L~~GA~~K~~VINMSFGNAALAYANLP
 DETKKAFDYAKSKGVSI~~V~~T~~S~~AGNDSSFGGK~~P~~RLPLADHPDYGVVGT~~P~~AAADSTLT~~V~~ASYS~~P~~D~~K~~QLTET
 ATVK~~T~~DDHQDKEMPV~~I~~STN~~R~~EP~~N~~KAYDYAYANRG~~T~~KEDDFKDVEGKIALIERGDIDFKDKIANAKKA
 GAVGVI~~I~~YDNQDKGF~~P~~IELPNVDQMPAAF~~I~~SR~~D~~GL~~L~~KDN~~P~~PKT~~I~~TFNATPKV~~L~~PTAS~~G~~TKLSRFSS
 30 WGLTADGN~~I~~KPDIAAPGQDILSSVANNKYAKLS~~G~~TSMSAPLVAGIM~~G~~LLQKQYETQY~~P~~DMTPSERLDL
 AKV~~L~~MSSAT~~A~~LYDEDEKAYFS~~P~~RQQGAGAVDAKKASAATMYVTDKDNTSSKVHLNNVSDKF~~E~~VT~~V~~T~~V~~
 HNKSDK~~P~~Q~~E~~LYQ~~V~~T~~V~~Q~~T~~D~~K~~V~~D~~G~~K~~H~~F~~AL~~A~~PK~~A~~LYETSWQ~~K~~IT~~I~~PANSS~~K~~Q~~V~~T~~V~~P~~I~~D~~A~~SR~~F~~SK~~D~~LLAQM
 KNGYFLEG~~V~~RFKQ~~D~~PT~~K~~KEELMS~~I~~PYIGFRGDFGN~~L~~SALEK~~P~~Y~~D~~SKDGSSYYHEANS~~D~~AKDQLDGDG
 35 LQFYALKNNFTALT~~T~~ESNPWT~~I~~IKAVKEGVENIEDIESSE~~E~~IT~~I~~FAGTF~~A~~KQ~~D~~DSHYYIHRHANGK
 PYAAISPNGDGNRDYVQFQGTFLRNAK~~N~~LVAE~~V~~LDKEGNVVWTSE~~V~~TEQVV~~V~~KN~~N~~NDLASTLGSTRFE
 KTRWDG~~K~~DKDGKV~~V~~ANGTYTYRV~~R~~Y~~T~~PISSG~~A~~KEQ~~H~~TD~~F~~D~~V~~IVDNTT~~P~~E~~V~~AT~~S~~AT~~F~~STEDSRL~~T~~LASK
 PKTSQPVY~~R~~ERIAYTYM~~D~~EL~~P~~T~~E~~Y~~I~~SPNEDG~~F~~TL~~P~~EEAETMEGATVPLKMSDFTYVVEDMAGNIT
 YTPVTKLLEGHSNKPEQDGSDQAPDKKPEAKPEQDGSGQTPD~~K~~KET~~K~~PEKDSSGQTPGKTPQKGQSS
 RTLEKRSSKRALATK~~A~~STRDQLPTTNDKDTNRLHLLKLVMTTFFLG

40 GBS 276 contains an N-terminal leader or signal sequence region which is indicated by the underlined sequence at the beginning of SEQ ID NO: 27 above. In one embodiment, one or more amino acids from the leader or signal sequence region of GBS 276 are removed. An example of such a GBS 276 fragment is set forth below as SEQ ID NO: 28.

SEQ ID NO: 28

5 QSDIKANTVTEDTPATEQAVEPPQPIAVSEESRSSKETKTSQTPSDVGETVADDANDLAPQAPAKTAD
 TPATSKATIRDLNDPSHVKTLOEKAGKGAGTVVAVIDAGFDKNHEAWRLTDKTKARYQSKENLEKAKK
 EHGITYGEWVNDKVAYYHDYSKDGKNAVDQEHGTHVSGILSGNAPSEMKEPYRLEGAMPEAQLLMLRV
 EIVNGLADYARNYAQAIRDAVNGLGAKVINMSFGNAALAYANLPDETKKAFDYAKSKGVSIANTSAGNDS
 10 SFGGKPRPLADHPDYGVVGT PAAADSTLTVASYSPDKQLTETATVKTDDHQDKEMPVISTNRFEPNK
 AYDYAYANRGTKEDDFKDVEGKIALIERGDIDFKDKIANAKKAGAVGVLIIYDNQDKGFPIELPNVDQM
 PAAFISRRDGLLLKDNPPTITFNATPKVLPTASGTLKLSRFSSWGLTADGNIKPDIAAPGQDILSSVA
 NNKYAKLSGTMSAPLVAGIMGLLQKQYETQYPDMTPSERLDLAKKVLMSATALYDEDEKAYFS PRQ
 QGAGAVDAKKASAATMYVTDKNTSSKVHLNNVSDKFEVTVTVHNKSDKPQELYQVTVQTDKVDGKH
 FALAPKALYETSWQKITIPANSSKQVTVPIADASRFSKDLLAQMKNGYFLEGFVRFQDPTKEELMSIP
 15 YIGFRGDFGNLSALEKPIYDSKDGSSYYHEANSDAKDQLDGDGLQFYALKNNFTALTTESNPWTIKA
 VKEGVENIEDIESSEITETIFAGTFAKQDDDSHYIYHRHANGKPYAAISPNGDGNRDYVQFQGTFRLN
 AKNLVAEVLDKEGNVWTSEVTEQVVKNYNNDLASTLGSTRFEKTRWDGKDKDGKVANGTYTYRVRY
 20 TPISSGAKEQHTDFDVIVDNTPEVATSATFSTEDSRLLTASLKPPTSQPVYERIAYTMDDELPTTE
 YISPNEDEGTFTLPEEAETMEGATVPLKMSDFTYVVEDMAGNITYTPVTKLLEGHSNKPEQDGSDQAPD
 KKPEAKPEQDGSGQTPDKKETKPEKDSSGQTPGKTPQKGQSSRTLEKRSSKRALATKASTRDQLPTT
 NDKDTNRLHLLKLVMTTFFLG

25 **GBS 276 contains a C-terminal transmembrane and/or cytoplasmic region which is indicated by the underlined sequence near the end of SEQ ID NO: 27 above. In one embodiment, one or more amino acids from the transmembrane or cytoplasmic regions of GBS 276 are removed. An example of such a GBS 276 fragment is set forth below as SEQ ID NO: 29.**

25

SEQ ID NO: 29

30 MRKKQKLPDFKLAIALISTSILLNAQSDIKANTVTEDTPATEQAVEPPQPIAVSEESRSSKETKTSQTPSDVGETVADDANDLAPQAPAKTADTPATSKATIRDLNDPSHVKTLOEKAGKGAGTVVAVIDAGFDKNHEAWRLTDKTKARYQSKENLEKAKKAPSEMKEPYRLEGAMPEAQLLMLRVIEVNGLADYARNYAQAIRDAVNGLGAKVINMSFGNAALAYANLPDETKKAFDYAKSKGVSIANTSAGNDSFGGKPRPLADHPDYGVVGT PAAADSTLTVASYSPDKQLTETATVKTDDHQDKEMPVISTNRFEPNKAYDYAYANRGTKEDDFKDVEGKIALIERGDIDFKDKIANAKKAGAVGVLIIYDNQDKGFPIELPNVDQMOMPAAFISRRDGLLLKDNPPTITFNATPKVLPTASGTLKLSRFSSWGLTADGNIKPDIAAPGQDILSSVANNKYAKLSGTMSAPLVAGIMGLLQKQYETQYPDMTPSERLDLAKKVLMSATALYDEDEKAYFS PRQGAGAVDAKKASAATMYVTDKNTSSKVHLNNVSDKFEVTVTVHNKSDKPQELYQVTVQTDKVDGKH FALAPKALYETSWQKITIPANSSKQVTVPIADASRFSKDLLAQMKNGYFLEGFVRFQDPTKEELMSIPYIGFRGDFGNLSALEKPIYDSKDGSSYYHEANSDAKDQLDGDGLQFYALKNNFTALTTESNPWTIKA VKEGVENIEDIESSEITETIFAGTFAKQDDDSHYIYHRHANGKPYAAISPNGDGNRDYVQFQGTFRLNAKVLVAEVLDKEGNVWTSEVTEQVVKNYNNDLASTLGSTRFETPISSGAKEQHTDFDVIVDNTPEVATSATFSTEDSRLLTASLPKPTSQPVYERIAYTMDDELPTTEYISPNEDEGTFTLPEEAETMEGATVPLKMSDFTYVVEDMAGNITYTPVTKLLEGHSNKPEQDGSDQAPDKEKDSSGQTPGKTPQKGQSSRTLEKRSSKRALATKASTRDQLPTTNDKDTNRLHLLKLVMTTFFLG

45 **In one embodiment, one or more amino acids from the leader or signal sequence region and one or more amino acids from the transmembrane or cytoplasmic regions of GBS 276 are removed. An example of such a GBS 276 fragment is set forth below as SEQ ID NO: 30.**

SEQ ID NO: 30

5 QSDIKANTVTEDTPATEQAVEPPQPIAVSEESRSSKETKTSQTPSDVGETVADDANDLAPQAPAKTAD
 TPATS KATIRDLNDPSHVKTLOEKAGKGAGTVVAVIDAGFDKNHEAWRLTDKTKARYQSKENLEKAKK
 EHGITYGEWVNDKVAYYHDYSKDGKNAVDQEHGTHVSGILSGNAPSEMKEPYRLEGAMPEAQLLLMRV
 EIVNGLADYARNYAQAIRDAVNLGAKVINMSFGNAALAYANLPDETAKSKGVSIVTSAGNDS
 10 SFGGKPRLPLADHPDYGVVGTAAADSTLTVASYSPDKQLTETATVKTDDHQDKEMPVISTNRFEPNK
 AYDYAYANRGTKEDDFKDVEGKIALIERGDIIFKDKIANAKKAGAVGLIYDNQDKGFPIELPNVDQM
 PAAFI SRRDGLLLKDNPBKTIIFNATPKVLPASGKLSRFSSWGLTADGNIKPDIAPGQDILSSVA
 NNKYAKLSGTSMSAPIVAGIMLLQKQYETQYPDMTPSERLDLAKKVLMSATALYDEDEKAYFSPRQ
 QGAGAVDAKKASAATMYVTDKDNTSSKVHLNNVSDKFEVTVTVHNKSDKPQELYQVTQTDKVDGKH
 15 FALAPKALYETSWQKITIPANSSQVTVPIADRSFSKDLLAQMKNGYFLEGFVRFQDPTKEELMSIP
 YIGFRGDFGNLSALEKPIYDSKDGSSYYHEANSDAKQLDGDGLQFYALKNNFTALTTESNPWTIIKA
 VKEGVENIEDIESSEITETIFAGTFAKQDDDSHYIYIHRHANGKPYAAISPNGDGNRDYVQFQGTFLRN
 AKNLVAEVLDKEGNVWTSEVTEQVVKNYNNLASTLGSTRFEKTRWDGKDKGKVVANGTYYRVRY
 TPISSGAKEQHTDFDVIDVNTPEVATSATFSTEDSRLLTASPKTSQPVYRERIAYTYMDEDLPTTE
 20 YISPNEGTFTLPEEAETMEGATVPLKMSDFTYVVEDMAGNITYTPVTKLLEGHSNKPEQDGSDQAPD
 KKPEAKPEQDGSGQTPKKETKPEKDSSGQTPGKTPQKGQSSRTLEKRSSKRALATK

Further description of GBS 276 can be found in the following references: Qi Chen et al., "Immunization with C5a Peptidase or Peptidase-Type III Polysaccharide conjugate Vaccines Enhances Clearance of Group B Streptococci from Lungs of Infected Mice", Infection and Immunity (2002) 70 (11):6409 – 6415; Beckmann et al., "Identification of Novel Adhesions from Group B Streptococci by Use of Phage Display Reveals that C5a Peptidase Mediates Fibronectin Binding" Infection and Immunity (2002) 70(6):2869 – 2876; Cheng et al., "The Group B Streptococcal C5a Peptidase Is Both a Specific Protease and an Invasin" Infection and Immunity (2002) 70(5) 2408 – 2413; and Cheng et al., "Antibody against Surface-Bound C5a Peptidase Is Opsonic and Initiates Macrophage Killing of Group B Streptococci" Infection and Immunity (2001) 69(4):2302 – 2308.

GBS 305

GBS 305 refers to a UDP-N-acetylmuramoylalanine--D-glutamate ligase, also referred to as Mur D. Nucleotide and amino acid sequences of GBS 305 sequenced from serotype V isolated strain 2603 V/R are set forth in Ref. 2 as SEQ ID 207 and SEQ ID 208. These sequences are set forth below as SEQ ID NOS 31 and 32:

SEQ ID NO. 31

35 ATGGGACGAGTAATGAAAACAATAACAAACATTGAAAATAAAAAGTTTAGTCCTGGTTAGCACG
 ATCTGGAGAAGCTGCTGCACGTTGTTAGCTAACGTTAGGAGCAATAGTGACAGTTAATGATGGCAAAC
 CATTGATGAAAATCCAACAGCACAGTCTTGTGGAAGAGGGTATTAAAGTGGTTGGTAGTCAT
 40 CCTTAGAATTGTTAGATGAGGATTTGTTACATGATTAAACCTCAGGAATACCTATAACAATCC
 TATGGTCAAAAAGCATTAGAAAACAATCCCTGTTGACTGAAGTGGATTAGCATACTTAGTT
 CAGAATCTCAGCTAATAGGTATTACAGGCTCTAACGGAAAACGACAACGACAACGATGATTGCAGAA
 GTCTTAAATGCTGGAGGTAGAGAGGTTGTTAGCTGGAAATATCGGCTTCTGCTAGTGAAGTTGT
 45 TCAGGCTGCGAATGATAAAGATACTCTAGTTATGGAATTATCAAGTTTCAGCTAATGGGAGTTAAGG
 AATTTCGTCCTCATATTGCAAGTAATTACTAATTAAATGCCAACTCATTAGATTATCATGGGTCTTT
 GAAGATTATGTTGCTGCAAAATGGAATATCCAAAATCAAATGTCTCATCTGATTGTTGGTACTTAA
 TTTAATCAAGGTATTCTAAAGAGTTAGCTAAAACACTAAAGCAACAATCGTCCTTCTACTA
 CGGAAAAGTTGATGGTCTACGTACAAGACAAGCAACTTTCTATAAAGGGGAGAATATTATGTCA

5 GTAGATGACATTGGTGTCCCAGGAAGCCATAACGTAGAGAATGCTCTAGCAACTATTGCGGTTGCTAA
 ACTGGCTGGTATCAGTAATCAAGTTATTAGAGAAACTTAAAGCAATTGGAGGTGTTAACACCGCT
 TGCAATCACCTCGGTAAAGGTTATGGTATTAGTTCTATAACGACAGCAAGTCAACTAATATATTGGCA
 ACTCAAAAAGCATTATCTGGCTTGTATAACTAAAGTTATCCTAATTGCAGGAGGTCTGATCGCG
 TAATGAGTTGATGAATTGATACAGATATCACTGGACTTAAACATATGGTTGTTAGGGGAATCGG
 10 CATCTCGAGTAAAACGTGCTGCACAAAAGCAGGAGTAACCTATAGCGATGCTTAGATGTTAGAGAT
 GCGGTACATAAAGCTTATGAGGTGGCACAACAGGGCGATGTTATCTGCTAAGTCCTGCAAATGCATC
 ATGGGACATGTATAAGAATTGAAAGTCCGTGGTATGAATTGATACTTCGAAAGTCTAGAG
 GAGAG

10 **SEQ ID NO. 32**
 15 MGRVMKTITT FENKKVLVLGLARSGEAAARLLAKLGAIVTVNDGKPFDENPTAQSLLEEGIKVVCGSH
 PLELLDEDFCYMIKNPGI PYNNPMVKKALEKQI PVLTEVELAYLVSESQLIGITGSNGKTTTTMIAE
 VLNAGGQRGLLAGNIGFPASEVVQAANDKDTLVMELSS FQLMGVKEFRPHIAVITNLMPHLDYHGSF
 EDYVAAKWNI QNQMSSSDFLVLFNQGISKELAKTTKATIVPFSTTEKVDGAYVQDKQLFYKGENIMS
 VDDIGVPGSHNVENALATIAVAKLAGISNQVIRETLSNFGGVKHRLQSLGVHGISFYNDSKSTNILA
TOKALSGFDNTKVILIAGGLDRGNEFDELI PDITGLKHMVVLGESASRVKRAAQKAGVTYS DALDVRD
AVHKAYEVAQQGDVILLSPANASWDMYKNFEVRGDEFIDTFESLRGE

20 GBS 305 contains an N-terminal leader or signal sequence region which is indicated by the underlined sequence at the beginning of SEQ ID NO: 32 above. In one embodiment, one or more amino acids from the leader or signal sequence region are removed from GBS 305. An example of such a GBS 305 fragment is set forth below as SEQ ID NO: 33.

25 **SEQ ID NO: 33**
 30 ITT FENKKVLVLGLARSGEAAARLLAKLGAIVTVNDGKPFDENPTAQSLLEEGIKVVCGSH PLELLDE
DFCYMIKNPGI PYNNPMVKKALEKQI PVLTEVELAYLVSESQLIGITGSNGKTTTTMIAE VLNAGGQ
RGLLAGNIGFPASEVVQAANDKDTLVMELSS FQLMGVKEFRPHIAVITNLMPHLDYHGS FEDYVAAK
 WNI QNQMSSS DFLVLFNQGISKELAKTTKATIVPFSTTEKVDGAYVQDKQLFYKGENIMS VDDIGVP
GSHNVENALATIAVAKLAGISNQVIRETLSNFGGVKHRLQSLGVHGISFYND SKSTNILA TOKALSG
FDNTKVILIAGGLDRGNEFDELI PDI TGLKHMVVLGESASRVKRAAQKAGVTYS DALDVRD AVHKAYE
 VAQQGDVILLSPANASWDMYKNFEVRGDEFIDTFESLRGE

35 GBS 305 contains a C-terminal transmembrane or cytoplasmic region indicated by the underlined sequence near the end of SEQ ID NO: 32 above. In one embodiment, one or more amino acids from the transmembrane or cytoplasmic regions are removed from GBS 305. An example of such a GBS 305 fragment is set forth below as SEQ ID NO: 34.

40 **SEQ ID NO: 34**
 45 MGRVMKTITT FENKKVLVLGLARSGEAAARLLAKLGAIVTVNDGKPFDENPTAQSLLEEGIKVVCGSH
 PLELLDEDFCYMIKNPGI PYNNPMVKKALEKQI PVLTEVELAYLVSESQLIGITGSNGKTTTTMIAE
 VLNAGGQRGLLAGNIGFPASEVVQAANDKDTLVMELSS FQLMGVKEFRPHIAVITNLMPHLDYHGSF
 EDYVAAKWNI QNQMSSSDFLVLFNQGISKELAKTTKATIVPFSTTEKVDGAYVQDKQLFYKGENIMS
 VDDIGVPGSHNVENALATIAVAKLAGISNQVIRETLSNFGGVKHRLQSLGVHGISFYNDSK

In one embodiment one or more amino acids from the leader or signal sequence region and one or more amino acids from the transmembrane or cytoplasmic regions are removed from GBS 305. An example of such a GBS 305 fragment is set forth below as SEQ ID NO: 35.

5 **SEQ ID NO: 35**

ITT FENKKVLVLGLARSGEAAARLLAKLGAIVTVNDGKPFDENPTAQSLLLEEGIKVVCGSHPLELLDE
 DFCYMIKNGPIPYNNPMVKKALEKQIPVLTEVELAYLVSESQQLIGITGSNGKTTTTMIAEVLNAGGQ
 RGLLAGNIGFPASEVVQAANDKDTLVMELSSFQLMGVKEFRPHIAVITNLMPHLDYHGSFEDYVAAK
 WNIQNQMSSDFLVLNFNQGISKELAKTTKATIVPFSTTEKVDGAYVQDKQLFYKGENIMSVDIGVP
 10 GSHNVENALATIAVAKIAGISNQVIRETLSNFGGVKHRLQSLGKVHGISFYNDSK

GBS 322

GBS 322 refers to a surface immunogenic protein, also referred to as "sip". Nucleotide and amino acid sequences of GBS 322 sequenced from serotype V isolated strain 2603 V/R are set forth in 15 Ref. 2 as SEQ ID 8539 and SEQ ID 8540. These sequences are set forth below as SEQ ID NOS 36 and 37:

SEQ ID NO. 36

ATGAATAAAAAGGTACTATTGACATCGACAATGGCAGCTCGCTATTATCAGTCGAAGTGTCAAGC
 20 ACAAGAAACAGATACGACGTGGACAGCACGTACTGTTTAGAGGTAAGGCTGATTGGTAAAGCAAG
 ACAAAATAATCATCATATACTGTGAAATATGGTGATACACTAACGTTATTCAGAAGCAATGTCAATT
 GATA TGAATGTCTTAGCAAAATAACATTGCAGATATCAATCTTATTATCCTGAGACAACACT
 GACAGTAACCTACGATCAGAAGAGTCATACTGCCACTTCAATGAAAATAGAAACACCAGCAACAAATG
 CTGCTGGTCAAACAACAGCTACTGTGGATTGAAAACCAATCAAGTTCTGTGCAGACCAAAAGTT
 25 TCTCTCAATACAATTCCGAAGGTATGACACCAAGCAGCAACAAACGATTGTTCGCAATGAAGAC
 ATAT TCTTCTGCGCCAGCTTGAAATCAAAGAAGTATTAGCACAAGAGCAAGCTGTTAGTCAGCAG
 CAGCTAATGAACAGGTATCACCAGCTCCTGTGAAGTCAGTACTTCAGAAGTTCCAGCAGCTAAAGAG
 GAAGTTAACCAACTCAGACGTCACTCAGTCAACAAACAGTATCACCAGCTTCTGTGCCGCTGA
 30 AACACCAAGCTCCAGTAGCTAAAGTAGCACCAGTAGCAGCTCCAGCAGTCCAGTGGCAAGTGTAA
 AAGTAGTCACTCCTAAAGTAGAAACTGGTGATCACCAGAGCATGTACAGCTCCAGCAGTCCCTGTG
 ACTACGACTTCACCAGCTACAGACAGTAAGTTACAAGCGACTGAAGTTAACAGCGTTCCGGTAGCACA
 AAAAGCTCCAACAGCAACACCGGTAGCACAACCAGCTTCACAAACAAATGCAGTAGCTGCACATCCTG
 AAAATGCAGGGCTCCAACCTCATGTTGCAGCTTATAAAGAAAAAGTAGCGTCAACTTATGGAGTTAAT
 35 GAATTCAGTACATACCGTGCAGGGAGATCCAGGTGATCATGGTAAAGGTTAGCAGTTGACTTTATTGT
 AGGTACTAATCAAGCACTGGTAATAAAGTTGCACAGTACTCTACACAAATATGGCAGCAAATAACA
 TTTCATATGTTATCTGGCAACAAAAGTTTACTCAAATACAAACAGTATTATGGACCTGCTAATACT
 TGGAAATGCAATGCCAGATCGTGGTGGCGTTACTGCCAACACTATGACCACGTTACGTATCATTTAA
 CAAAATATATAAAAAGGAAGCTATTGGCTTTTATATGCCCTGAATAGACTTCAAGGTTCT
 40 TATAATTTTATTA

SEQ ID NO. 37

MNKKVLLTSTMASLLSVASVQAQETDTWTARTVSEVKADLVQDNKSSYTVKYGDTLSVISEAMSI
 DMNVIAKINNIADINLIYPETTLTVTYDQKSHTATSMKIEPATNAAGQTTATVDLKTNQVSADQKV
 45 SLNTISEGMTPEAATTIVSPMKTYSAPALKSKEVLAQEQAQSAAANEQVSPAPVKSITSEVPAKE
 EVKPTQTSVSQSTTVSPASVAAETPAPVAKVAPVRTVAAPRVASVTVPKVETGASPEHVSAPAVPV
 TTSPATDSKLQATEVKSVPVAQKAPTPVAPQPASTTNAVAAHPENAGLQPHVAAYKEKVASTYGVN
 EFSTYRAGDPGDHGKGLAVDFIVGTNQALGNKVAQYSTQNMAANNISYVIWQQKFYSNTNSIYGPANT
 WNAMPDRGGVTANHYDHVHSFNK

GBS 322 contains an N-terminal leader or signal sequence region which is indicated by the underlined sequence near the beginning of SEQ ID NO: 37. In one embodiment, one or more amino acids from the leader or signal sequence region of GBS 322 are removed. An example of such a GBS 5 322 fragment is set forth below as SEQ ID NO: 38.

SEQ ID NO: 38

DLVKQDNKSSYTVKYGDTLSVISEAMSI DMNVLA KINNIADINLIYPETTLTVYDQKSHTATSMKIE
 10 TPATNAAGQTTATVDLKTNQVSVA DQKVS LNTISEG MTP EAATTIVSPMKT YSSA PALKSKEVLAQEQ
 AVSQAAANEQVS PAPVKSITSEVPAKEEVKPTQTSVQSTTVSPASVAAETPAPVAKVAPV RVTVAAP
 RVASVKVVTPKVETGASPEHVSAPAVPVTTSPATDSKLQATEVKSV PVAQKAPTATPVAQ PASTTN
 VAAHPENAGLQPHVAAYKEKVASTYGVNEFSTYRAGDPGDHGKGLAVDFIVGTNQALGNKVAQYSTQN
 MAANNISYVIWQQKFYSNTNSIYGPANTWNAMPDRGGVTANHYDHVHVSFNK

15 **GBS 330**

GBS 330 refers to a pyruvate kinase, also referred to as "pyk". Nucleotide and amino acid sequences of GBS 330 sequenced from serotype V isolated strain 2603 V/R are set forth in Ref. 2 as SEQ ID 8791 and SEQ ID 8792. These sequences are set forth below as SEQ ID NOS 39 and 40:

20 **SEQ ID NO. 39**

ATGAATAAACCGTAAAAATCGTTGCAACACTGGTCCTCGGGTTGAATTCCGTGGTGGTAAGAAGTT
 TGGTGAGTCTGGATACTGGGGTGAAGGCCCTGACGTAGAACGCTTCAGCAGAAAAATTGCTCAATTGA
 TTAAAGAAGGTGCTAACGTTTCCGTTCAACTCTCACATGGAGATCATGCTGAGCAAGGAGCTCGT
 ATGGCTACTGTTCGTAAAGCAGAACAGAGATTGCAGGACAAAAAGTTGGCTCCCTTGATACTAAAGG
 25 ACCTGAAATTCTGACAGAACCTTTGAAGATGGCAGATTCCATTCAATACAAACAGGTACAAAAT
 TACGTGTTGCTACTAACAGCAAGGTATCAAATCAACTCCAGAACGTGATTGCATTGAATGTTGCTGGGA
 CTTGACATCTTGATGACGTTGAAGTTGGTAAGCAAATCCTGTTGATGATGGTAAACTAGGTCTTAC
 TGTGTTGCAAAAGATAAACAGACACTCGTGAATTGAAGTAGTTGTTGAGAATGATGGCCTTATTGGTA
 30 AACAAAAAGGTGTAAACATCCCTTATAC TAAATCCCTTCCCAGCACTTGCAGAACCGATAATGCT
 GATATCCGTTTGGACTTGAGCAAGGACTTAACCTTATTGCTATCTCATTGTA CGTACTGCTAAAGA
 TGTTAATGAAGTTCGTGTATTGTGAAGAAACTGGSMATGGACACGTTAAGTTGTTGCTAAATG
 AAAATCAACAAGGTATCGATAATATTGATGAGATTATCGAAGCAGCAGATGGTATTATGATTGCTCGT
 GGTGATATGGGTATCGAAGTCCATTGAAATGGTCCAGTTACAAAAATGATCATTACTAAAGT
 35 TAATGCAGCTGGTAAAGCAGTTATTACAGCAACAAATATGCTTGAACAAATGACTGATAAACCAACCGTG
 CGACTCGTCAGAAGTATCTGATGTCTCAATGCTGTTATTGATGGTACTGATGCTACAATGCTTCA
 GGTGAGTCAGCTAATGGTAAATAACCCAGTTGAGTCAGTTCGTACAATGGCTACTATTGATAAAAATGC
 TCAAACATTA CACTCAATGAGTATGGTCCCTAGACTCATCTGCATTCCACGTAATAACAAAATGATG
 TTATTGCATCTGC GGTTAAAGATGCAACACACTCAATGGATATCAAACATTGTTGAACAATTACTGAA
 40 ACAGGTAATACAGCTCGTGC CATTCTAAATTCCGTCAGATGCAGACATTGGCTGTTACATTG
 TGAAAAAGTACAACGTTATTGATGATTA ACTGGGTGTATCCCTGCTTGCAGACAAACCAGCAT
 CTACAGATGATATGTTGAGGTTGCAGAACGTGTAGCATTGAAGCAGGATTGTTGAATCAGGGCAG
 AATATCGTTATCGTGCAGGTGTTCTGTAGGTACAGGTGGAACTAACACAATGCGTGTACTGT
 TAAA

SEQ ID NO. 40

5 MNKRVKIVATLGPRAVEFRGGKKFGESGYWGESLDVEASAECIAQLIKEGANVFRFNFSHGDHAEQGAR
 MATVRKAEEIAGQKVGFLLDTKPEIRTELFDGADFHSYTTGKLRVATKQGIKSTPEVIALNVAGG
 LDI FDDVEVGKQILVDDGKLGTVFAKDKDTRFEVVENDGLIGKQKGVNIPYTKIPFPALAERDNA
 DIRFGLEQGLNFIAISFVRTAKDVNEVRAICEETGXGHVKLFAKIENQQGIDNIDEIIEAADGIMIAR
 GDMGIEVPFEMVPVYQKMIITKVNAAGKAVITATNMLETMDKPRATRSEVSDFNAVIDGTDATMLS
 10 GESANGKYPVESVRTMATIDKNAQTLLNEYGRLDSSAFPRNNKTDVIASAVKDATHSMDIKLVVTITE
 TGNTARAISKFRPDADILAVTFDEKVQRSLMINWGVIPVLADKPASTDDMFEVAERVALEAGFVESGD
 NIVIVAGVPVGTGGTNTMRVRTVK

GBS 338

GBS 338 refers to a Sat D protein. Nucleotide and amino acid sequences of GBS 338
 sequenced from serotype V isolated strain 2603 V/R are set forth in Ref. 2 as SEQ ID 8637 and SEQ
 15 ID 8638. These sequences are set forth below as SEQ ID NOS 41 and 42:

SEQ ID NO. 41

20 TTGTCTGCTATAATAGACAAAAAGGTGGTGATATTATGTATTTAGCATTAAATCGGTGATATCATTAA
 TTCAAAACAGATACTTGAACGTGAAACTTCAACAGTCTTCAGCAACTAATGACCGAACTATCTG
 ATGTATATGGTGAAGAGCTGATTCTCATTCACTATTACAGCTGGTATGAATTCAAGCTTATTG
 AAACCATCAAAAAGGTATTCAAATTATTGACCATTCAACTAGCTCTAAACCTGTTAATGTAAG
 GTTCGGCCTCGGTACAGGAAACATTATAACATCCATCAATTCAAATGAAAGTATCGGTGCTGATGGTC
 25 CTGCCTACTGGCATGCTCGCTCAGCTATTAAATCATATAACATGATAAAATGATTATGGAACAGTTCAA
 GTAGCTATTGCGCTTGATGATGAAGACCAAAACCTGAAATTAAACACTAAATAGCTCATTCAAGCTGG
 TGATTTATCAAGTCAAATGGACTACAAACCTTTCAAATGCTTGAGCACTTAATACTCAAGATA
 ATTATCAAGAACAAATTCAACATCAAAGTTAGCCCACTGGAAATATTGAACCTAGTGCCTGACT
 30 AACGCCTTAAAGCAAGCGGTCTGAAGATTACTTAAGAACGAGAACACAGGCAGCCGATCTATTAGT
 TAAAAGTTGCACTCAAACAAAGGGGGAAAGCTATGATTTC

SEQ ID NO. 42

35 MSAIIDKKVVIFMYLALIGDIINSKQILERETFQQS FQQLMTELSDVYGEELISPFTITAGDEFQALL
 KPSKKVFQIIDHQLALKPVNVRFLGTTGNIITSINSNESIGADGPAYWHARSAINHIHDKN DYGTVO
 VAICLDDEDQNL ELTLNSLISAGDFIKSKWTTNHFQMLEHLILQDNYQEQQHQKLAQLENIEPSALT
 KRLKASGLKIYLRTRTQAADLLVKSCQTKGGSYDF

40 GBS 338 may contain an N-terminal leader or signal sequence region which is indicated by
 the underlined sequence at the beginning of SEQ ID NO: 42 above. In one embodiment, one or more
 amino acids from the leader or signal sequence region are removed from GBS 338. An example of
 such a GBS 338 fragment is set forth below as SEQ ID NO: 43.

40

SEQ ID NO: 43

45 MYLALIGDIINSKQILERETFQQS FQQLMTELSDVYGEELISPFTITAGDEFQALLKPSKKVFQIIDH
 IQLALKPVNVRFLGTTGNIITSINSNESIGADGPAYWHARSAINHIHDKN DYGTVOVAICLDDEDQNL
 ELTLNSLISAGDFIKSKWTTNHFQMLEHLILQDNYQEQQHQKLAQLENIEPSALT KRLKASGLKIYL
 RTRTQAADLLVKSCQTKGGSYDF

GBS 361

GBS 361 refers to a cylI protein. Nucleotide and amino acid sequences of GBS 361 sequenced from serotype V isolated strain 2603 V/R are set forth in Ref. 2 as SEQ ID 8769 and SEQ ID 8770. These sequences are set forth below as SEQ ID NOS 44 and 45:

5

SEQ ID NO. 44

ATGAGCGTATATGTTAGTGGAAAGGAATTATTCTTCTTGGAAAGAATTATAGCGAGCATAAACA
 GCATCTCTCGACTAAAAGAAGGAATTCTAAACATTATATAAAAATCACGACTCTATTAGAAT
 CTTATACAGGAAGCATAACTAGTGACCCAGAGGTTCTGAGCAATACAAAGATGAGACACGTAATTT
 10 AAATTTGCTTTACCGCTTGAAGAGGCTCTGCTTCTCAGGTGTTAATTAAAAGCTTATCATAA
 TATTGCTGTGTTAGGGACCTCACTGGGGAAAGAGTGCTGGTAAAATGCCTGTATCAATTG
 AAGAAGGAGAGCGTCAAGTAGATGCTAGTTATTAGAAAAGCATCTGTTACCATATTGCTGATGAA
 TTGATGGCTATCATGATATTGTGGGAGCTCGTATGTTATTCAACCGCCTGTTCTGCAAGTAATAA
 15 TGCCGTAATATTAGGAACACAATTACTCAAGATGGCATTGTGATTAGCTATTGTGGTGGCTGTG
 ATGAGTTAAGTGTATTTCTTAGCAGGCTTCACATCACTAGGAGCTTAATACAGAAATGGCATGT
 CAGCCCTATTCTGGAAAAGGAATCAATTGGGTGAGGGCGCTGGTTGTTGTTCTGCAAAGA
 TCAGTCCTAGCTAAATATGGAAAATTATCGGTGGTCTTATTACTCAGATGGTTATCATATAACAG
 CACCTAACGCAACAGGTGAAGGGCGGCACAGATTGCAAAGCAGCTAGTGACTCAAGCAGGTATTGAC
 TACAGTGAGATTGACTATATTACGGTCACGGTACAGGTACTCAAGCTAATGATAAAATGGAAAAAA
 20 TATGTATGGTAAGTTTCCCACACGACATTGATCAGCAGTACCAAGGGCAAACGGGTCAACTC
 TAGGGGCTGCAGGTATTATCGAATTGATTAATTGTTAGCGGCAATAGAGGAACAGACTGTACAGCA
 ACTAAAAATGAGATTGGATAGAAGGTTCCAGAAAATTGTCTATCATCAAAGAGAGAAATACCC
 AATAAGAAATGCTTAAATTTCGTTGCTTTGGAAATAATAGTGGTGTCTTGTCTTCTGCT
 25 TAGATTCACCTCTAGAACATTACCTGCTAGAGAAAATCTAAAATGGCTATCTTATCATCTGTTGCT
 TCCATTCTAAGAATGAATCACCTCTATAACCTATGAAAAGTTGCTAGTAATTCAACGACTTTGA
 AGCATTACGCTTAAAGGGCTAGACCACCCAAAATGTCAACCCAGCACAAATTAGAAAATGGATG
 ATTTTCCAAATGGTGCCTAACACAGCTCAAGCACTAATAGAAAGCAATATTAACTAAAAAAA
 CAAGATACTCAAAAGTAGGAATTGTATTACAACACTTCTGGACCAGTTGAGGTGTTGAAGGTAT
 TGAAAAGCAAATCACACAGAAGGATATGCACATGTTCTGCTCACGATTCCGTTACAGTAATGA
 30 ATGCAGCAGCTGGTATGTTCTATCATTTAAAATACAGGTCTTATCTGTATTCGACAAAT
 AGTGGAGCGCTTGATGGTATACAATATGCCAAGGAATGATGCGTAACGATAATCTAGACTATGTGAT
 TCTTGTCTGCTAATCAGTGGACAGACATGAGTTATGTGGCAACAATTAAACTATGATAGTC
 AAATGTTGTCGGTCTGATTATTGTTCAAGCACAAGTCCCTCTCGTCAAGCATTGGATAATTCCCT
 ATAATATTAGGTAGTAAACAATTAAAATAGCATAAAACATTACAGATGTGACTATTGTA
 35 TGCTGCGCTTCAAAATTATTACAGACTTAGGACTAACATAAAAGATATCAAAGGTTGCTTGG
 ATGAGCGGAAGAAGGCAGTTAGTCAGATTATGATTCTAGCGAATTGCTGAGTATTATAATATG
 CCAAACCTGCTCTGGCAGTTGGATTTCATCTAATGGTCTGGTAAGAACTGGACTACTGT
 TAATGAAAGTATAGAAAAGGCATTATTAGTCCTATCTTATTGATCTCGGTGGTATCTTTG
 CTATTATTGAAAAAAGG

40

SEQ ID NO. 45

MSVYVSGIGIISSLGKNYSEHKQHLFDLKEGISKHLYKNHDSILESYTGSITSDPEVPEQYKDETRNF
 KFAFTAFEEALASSGVNLKAYHNIAVCLGTSLLGKSAGQNALYQFEEGERQVDASLLEKASVYHIADE
 LMAYHDIVGASYVISTACSASNNAVILGTQLLQDGDCDLAICGGCDELSDISLAGFTSLGAINTEMAC
 45 QPYSSGKGINLGEAGFVVLVDQSLAKYGKIIIGLITS

DGYHITAPKPTGEGAAQIAKQLVTQAGID
 YSEIDYINGHGTGQANDKMEKNMYGKFPTTLLISSTKGQTGHTLGAAGIIELINCLAAIEEQTVPA
 TKNEIGIEGFENFVYHQKREYPIRNALNFSFAFGGNNSGVLLSSLDSPLETLPARENLKMAILSSVA
 SISKNESLSITYEVASNFNDFEALRFKGARPPKTVNPAQFRKMDDFSKMVAVTTAQALIESNINLKK
 QDTSKVGVIVFTTLSGPVEVVEGIEKQITTEGYAHVSASRFPFTVMNAAAGMLSIIFKITGPLSVISTN
 50 SGALDGIQYAKEMMRNDNLVILVSANQWTDMFSMWWQQLNYDSQMFVGSDYCSAQVLSRQALDNP

I I LGSKQLKYSHKTFTDVMT I FDAALQNLSDLGLTIKDIKGFVWNERKKAVSSDYDFLANLSEYYNM
PNLASQQFGFSSNGAGEELDYTVNESIEKGYYLVLSSYI FGGISFAIIIEKR

GBS 361 may contain an N-terminal leader or signal sequence region which is indicated by the underlined sequence at the beginning of SEQ ID NO: 45 above. In one embodiment, one or more 5 amino acids from the leader or signal sequence region are removed from GBS 361. An example of such a GBS 361 fragment is set forth below as SEQ ID NO: 46.

SEQ ID NO: 46

VSGIGIISLGKNYSEHKQHLFDLKEGISKHLYKNHDSILESYTGSITS DPEVPEQYKDETRNFKFAF
10 TAFEEALASSGVNLKAYHNIAVCLGTSLLGGKSAGQNLYQFEEGERQVDASLLEKASVYHIADELMAY
HDIVGASYVISTACSASNNAVILGTQLLQDGDCDLAICGGCDELSDISLAGFTSLGAINTEMACQPPYS
SGKGINLGEAGAGFVVLVKDQSLAKYGTIGGLITSDGYHITAPKPTGEGAAQIAKQLVTQAGIDYSEI
DYINGHGTGTQANDKMEKNMYKFFPTTLISSTKGQTGHTLGAAGIIELINCLAAIEEQTVPATKNE
15 IGIEGFPEVFVYHQKREYPIRNALNFSFAFGGNNSGVLLSSLDSPLETLPARENLKMAILSSVASISK
KVGIVFTTLSGPVEVVEGIEKQITTEGYAHVSASRFPTVMNAAAGMLSIIFKITGPLSVISTNSGAL
DGIQYAKEMMRNDNLDYVILVSANQWTDMSFMWWQQLNYDSQMFVGSDYCSAQVLSRQALDNSPILG
SKQLKYSHKTFTDVMT I FDAALQNLSDLGLTIKDIKGFVWNERKKAVSSDYDFLANLSEYYNMPNLA
20 SGQFGFSSNGAGEELDYTVNESIEKGYYLVLSSYI FGGISFAIIIEKR

GBS 404

Nucleotide and amino acid sequences of GBS 404 sequenced from serotype V isolated strain 2603 V/R are set forth in Ref. 2 as SEQ ID 8799 and SEQ ID 8800. These sequences are set forth below as SEQ ID NOS 47 and 48:

25

SEQ ID NO. 47

ATGAAAATAGATGACCTAAGAAAAAGCGACAAATGTTGAAGATCGTCGCTCCAGTAGCGGAGGTTCAATT
CTCTAGCGGAGGAAGTGGATTACCGATTCTCAACTTTATTGCTGCGAGGGAGTTGGAAAACCAAGC
30 TTGTGGTTTTAATCATCTTACTGCTACTTGGCGGAGGGGACTAACAGCATTTAATGACTCATCC
TCACCTCTAGTTACCAATCTCAGAATGTCTCACGTTCTGTTGATAATAGCGCAACGAGAGAACAAAT
CGATTCGTTAATAAAAGTCCTGGCTCAACTGAGGATTCTGGTCACAAGAATTCCAACCCAAAGGTT
TTGGAAATTATAAGGAACCAAAACTTGGTCTTACACCAATTCAATTCAAACAGGTTGGTATAGGT
GAATCTGCTTCAGGACCATTATTGTTACAGCAGATAAAAAATCTATCTTGATATTCTTTACAA
TGAATTATCACATAATATGGTCTACTGGTATTGCTATGGCCTACGTACGCCACGAAGTTG
35 GTCACACATTCAAACAGAGTTAGGCATTATGGATAAGTATAATAGAATGCACACGGACTTACTAAG
AAAGAAGCAAATGCTTAAATGTTGGCTAGAACCTCAAGCAGATTATTATGCAGGGGTATGGCTCA
CTACATCAGGGGAAAAATCTCTAGAACAAAGGAGACTTGAAGAGGCCATGAATGCTGCCACGCCG
TCGGAGACGATAACCTTCAGAAAAGAAACCTACGGAAAATTAGTGCCTGATAGCTTACCCATGGAACA
40 GCTGAACAACGCCAACGTTGGTTAACAAAGGCTTCAATATGGTACATCCAACACGGTACATT
CTCCGTAGAACATCTA

SEQ ID NO. 48

MKIDDLRKSDNVEDRRSSSGGSFSSGGSGLPILQLLLRGSWTKLVVLI I LLLLGGGLTSIFNDSS
SPSSYQSQNVSRSVDNSATREQIDFVNKVLGSTDWFWSQE FQTQGFGNYKEPKLVLYTNSIQTGCGIG
45 ESASGPFYCSADKKIYLDISFYNELSHKYGATGDFAMAYVIAHEVGHHIQTELGIMDKYNRMRHGLTK
KEANALNVRLELQADYYAGVWAHYIRGKNLLEQGDFEEAMNAAHAVGDDTLQKETYGKLPDSFTHGT
AEQRQRWFNKGQYGDIQHGDTSVEHL

GBS 690

Nucleotide and amino acid sequences of GBS 690 sequenced from serotype V isolated strain 2603 V/R are set forth in Ref. 2 as SEQ ID 9965 and SEQ ID 9966. These sequences are set forth as 5 SEQ ID NOS 49 and 50 below:

SEQ ID NO. 49

ATGAGTAAACGACAAATTAGGAATTAGTAAAAAAGGAGCAATTATATCAGGGCTCTCAGTGGCACT
 10 AATTGTAGTAATAGTGGCTTTATGGGTACAATCTCAACCTAATAAGAGTCAGTAAAAACTAACT
 ACAAAGTTTAATGTTAGAGAAGGAAGTGTTCGTCCTCAACTCTTTGACAGGAAAGCTAAGGCT
 AATCAAGAACAGTATGTGTATTGATGCTAATAAAGGTAAATCGAGCAACTGTCACAGTTAAAGTGGG
 15 TGATAAAAATCACAGCTGGTCAGCAGTTAGTCATATGATACAACAACAGCAGCCTACGACA
 CTGCTAATCGTCATAATAAAGTAGCGCGTCAGATTAATAATCTAAAGACAAACAGGAAGTCTTCCA
 GCTATGGAATCAAGTGTCAATCTTCATCATCACAAGGACAAGGGACTCAATCGACTAGTGGTGC
 GACGAATCGTCTACAGCAAAATTATCAAAGTCAAGCTAATGCTTCATACAAACCAACAATTCAAGATT
 20 TGAATGATGCTTATGCGAGATGCACAGGAGAAGTAAATAAAGCACAAAAAGCATTGAATGATACTGTT
 ATTACAAGTGACGTATCAGGGACAGTTGTAAGTTAATAGTGTATTGATCAGCTTCAGCTCAAAACTAG
 TCAAGTACTTGTCCATGTAGCAACTGAAGGTAACACTCAAGTACAAGGAACGATGAGTGTGATT
 25 TGGCTAATGTTAAAAAAGACCAGGCTTTAAATAATCTAAGGTCTATCCTGACAAGGAATGGAA
 GGTAAAATTCTATATCTCAAATTATCCAGAAGCAGAAACAACAAATGACTCTAATAACGGCTC
 TAGTGCTGTTAAATTATAAATAAAGTAGATATTACTAGCCCTCTCGATGCATTAAACAAGGTTTA
 CCGTATCAGTTGAAGTAGTTAATGGAGATAAGCACCTTATTGTCCTACAAGTCTGTGATAAACAAA
 GATAATAAACACTTGTGTTGGGTATACAATGATTCTAATCGTAAATTCTCAAAGTTGAAGTCAAAT
 TGGTAAAGCTGATGCTAAGACACAAGAAATTTCAGGTTGAAAGCAGGACAAATCGTGGTTACTA
 35 ATCCAAGTAAACCTCAAGGATGGGAAAAATTGATAATATTGAATCAATCGATCTAACTCTAAT
 AAGAAATCAGAGGTGAAA

SEQ ID NO. 50

MSKRQNLGISKKGAIISGLSVALIVVIGGFLWVQSPNKSAVKTNYKVFNVREGSVSSSTLLTGKAKA
 30 NQEQQYVYFDANKGNRATVTVKVGDKITAGQQQLVQYDTTTAQAAAYDTANRQLNKVARQINNLKTTGSLP
 AMESSDQSSSSQGQGTQSTSGATNRQLQQNYQSQANASYNQQLQDLNDAYADAQAEVNKAQKALNDTV
 ITSDVSGTVVEVNSDIDPASKTSQVLVHVATEGKLQVQGTMSEYDLANVKKDQAVKIKSKVYPDKEWE
 GKISYISNYPEAEANNNDNSNNGSSAVNYKYKVDITSPLDALKQGFTVSVEVNGDKHLIVPTSSVINK
 DNKHFVWVYNDSNRKISKVEVKIGKADAKTQEILSGLKAGQIVVTNPSKTFKDQKIDNIESIDLNSN
 35 KKSEVK

GBS 690 contains an N-terminal leader or signal sequence region which is indicated by the underlined sequence at the beginning of SEQ ID NO: 50 above. In one embodiment, one or more amino acids from the leader or signal sequence region of GBS 690 are removed. An example of such 40 a GBS 690 fragment is set forth below as SEQ ID NO: 51.

SEQ ID NO: 51

FLWVQSPNKSAVKTNYKVFNVREGSVSSSTLLTGKAKANQEQQYVYFDANKGNRATVTVKVGDKITAG
 45 QQQLVQYDTTTAQAAAYDTANRQLNKVARQINNLKTTGSLPAMESSDQSSSSQGQGTQSTSGATNRQLQQ
 NYQSQANASYNQQLQDLNDAYADAQAEVNKAQKALNDTVITSDVSGTVVEVNSDIDPASKTSQVLVHV
 ATEGKLQVQGTMSEYDLANVKKDQAVKIKSKVYPDKEWE
 GKISYISNYPEAEANNNDNSNNGSSAVNYK
 YKVDITSPLDALKQGFTVSVEVNGDKHLIVPTSSVINKDNKHFVWVYNDSNRKISKVEVKIGKADAK
 TQEILSGLKAGQIVVTNPSKTFKDQKIDNIESIDLNSNKKSEVK

GBS 691

GBS 691 refers to an iron compound ABC transporter, or a substrate binding protein.

Nucleotide and amino acid sequences of GBS 691 sequenced from serotype V isolated strain 2603

5 V/R are set forth in Ref. 2 as SEQ ID 3691 and SEQ ID 3692. These sequences are set forth as SEQ ID NOS 52 and 53 below:

SEQ ID NO. 52

ATGAA~~AAA~~ATTGGAATTATTGCTCACACTACTGACCTTCTTTGGTATCTGCGGACAACAAAC
 10 TAAACAAGAAAGCACTAAAACA~~T~~TTCTAAAATGCCTAAAATTGAAGGCTTCACCTATTATGGAA
 AAATT~~C~~CTGAAAATCCGAAAAAAGTAATTAATT~~T~~ACATATTCTTACACTGGGTATTATTAAA~~A~~CTA
 GGTGTTAATGTTCAAGTTACAGTTAGACTTAGAAAAAGATAGCCCCGTTTGGTAAACAACTGAA
 AGAAGCTAAAATTA~~A~~CTGCTGATGATACAGAAGCTATTGCCGCACAAAACCTGATTAA~~T~~CATGG
 15 TTTGATCAAGATCAAACATCAATACTCTGAAAAAAATTGCACCAACTT~~A~~GGTATTAAATATGGT
 GCACAAAATTATTAGATATGATGCCAGCCTGGGGAAAGTATTGGTAAAGAAAAGAAGCTAATCA
 GTGGGTAGCCAATGGAAA~~A~~CTCTCGCTGTCAAAAAGATTACACC~~A~~TCTTAAAGCCTA
 ACAC~~T~~ACTTTACTATTATGGATT~~T~~TATGATAAAAATATCTATT~~T~~ATATGGTAA~~T~~ATTGGACGC
 GGTGG~~A~~GA~~A~~CTAATCTATGATTCACTAGGTTATGCTGCCAGAAAAAGTCAAAAAGATGTCTTAA
 AAAAGGGTGGTTACCGTTCGCAAGAAGCA~~T~~CGGTGATTACGTTGGAGATTATGCCCTGTTAATA
 20 TAAACAAA~~A~~CGACTAAAAGCAGCTTCATC~~A~~CTAAAGAAAGT~~G~~ATG~~T~~CTGGAAGAATTACAGCT
 GTCAAAAAGGGCACATCATAGAAAGTA~~A~~CTACGACGT~~T~~TTATTCTCTGACCCT~~T~~ATCTTAGA
 AGCTCAATTAAAATCATTTACAAAGGCTATCAAAGAAAATACAAAT

SEQ ID NO. 53

MKKIGTIVLLLTFFLVSCGQQTKQESTKTTISKMPKIEGFTYYGKIPENPKVINFTYSYTGYLLKL
 25 GVNVSSYSLDLEKDSPVFGKQLKEAKKLTADDTEAIAAAQPKDLIMVFDQDPNINTLKKIAPTLVIKYG
 AQNYLDMMMPALGKVFGKEKEANQWVSQWKT~~K~~TLAVKKDLHHILKPNTTFTIMDFYDKNIYLYGNNFGR
 GGELI~~Y~~DSLGYAAPEKVKKDVFKKGWFTVSQEAIGDYVGDYALVNINKTKAASSL~~K~~ESDVWKNLPA
 VKKGHI~~I~~ESNYDVFYFSDPLSLEAQLKSFTKAIKENTN

30 GBS 691 contains an N-terminal leader or signal sequence region which is indicated by the underlined sequence at the beginning of SEQ ID NO: 53 above. In one embodiment, one or more amino acids are removed from the leader or signal sequence region of GBS 691. An example of such a GBS 691 fragment is set forth below as SEQ ID NO: 54.

SEQ ID NO: 54

EGFTYYGKIPENPKV~~INFTYSYTGYLLKLGVNVSSYSL~~DLEKDSPVFGKQLKEAKKLTADDTEAIAAAQPKDLIMVFDQDPNINTLKKIAPTLVIKYGAQNYLDMMMPALGKVFGKEKEANQWVSQWKT~~K~~TLAVKKDLHHILKPNTTFTIMDFYDKNIYLYGNNFGRGGELI~~Y~~DSLGYAAPEKVKKDVFKKGWFTVSQEAIGDYVGDYALVNINKTKAASSL~~K~~ESDVWKNLPA

40 GBS 691 contains a C-terminal transmembrane or cytosolic region which is indicated by the underlined sequence at the end of SEQ ID NO: 53 above. In one embodiment, one or more amino acids are removed from the transmembrane or cytosolic region of GBS 691. An example of such a GBS 691 fragment is set forth below as SEQ ID NO: 55.

SEQ ID NO: 55

MKKI GI IVLTLTFLVSCGQQTKQESTKTTISKMPKIEGFTYYGKIPENPKVINFYSYTGYLLKL
 GVNVSYSLDLEKDS PVFGKQLKEAKKL TADDTEAIAAQKPDLIMVFDQDPNINTLKKIAPTLVIKYG
 5 AQNYILDMMMPALGKVFKEKEANQWVSQLKTKTLAVKKDLHHILKPNTTFTIMDFYDKNIYLYGNNFGR
 GGELIYDSIIGYAAPEKVKKDVFKKGWFTVSQEAIGDYVGDYALVNINKTTKAASSLKESDVWKNLPA
 VKKGHIIIESNYDVFYFSDPLSLEAQLKSFT

In one embodiment, one or more amino acids from the leader or signal sequence region and one or more amino acids from the transmembrane or cytosplasmic region are removed from GBS 691.

10 One example of such a GBS 691 fragment is set forth below as SEQ ID NO: 56

SEQ ID NO: 56

EGFTYYGKIPENPKVINFYSYTGYLLKLGVNVSSYSLDLEKDS PVFGKQLKEAKKL TADDTEAIAA
 15 QKPDLIMVFDQDPNINTLKKIAPTLVIKYGAQNYILDMMMPALGKVFKEKEANQWVSQLKTKTLAVKKD
 LHHILKPNTTFTIMDFYDKNIYLYGNNFGRGGELIYDSIIGYAAPEKVKKDVFKKGWFTVSQEAIGDYV
 GDYALVNINKTTKAASSLKESDVWKNLPAVKKHIIIESNYDVFYFSDPLSLEAQLKSFT

Additional examples of GBS antigens which may be used in combination with GBS 80 are set forth below.

20 **GBS 4**

GBS 4 refers to another putative cell wall surface anchor family protein. Nucleotide and amino acid sequences of GBS 4 sequenced from serotype V isolated strain 2603 V/R are set forth in Ref. 2 as SEQ ID 1 and SEQ ID 2. These sequences are also set forth below as SEQ ID NOS 57 and 58:

25

SEQ ID NO. 57

ATGAAAGTGAAAATAAGATTAAACGATGGTAGCACTTACTGTCTAACATGTGCTACTTATTCACT
 AATCGGTTATGCTGATAACAAGTATAAGAAATACGTACACAGAGTGTGACTACGACCTATCTGAGG
 30 AGAAAAGATCAGATGAACTAGACCAAGTCTAGTACTGGTTCTCTCTGAAAATGAATCGAGTTCATCA
 AGTGAACCAGAAACAAATCCGTCAACTAATCCACCTACAACAGAACCATCGCAACCCCTCACCTAGTGA
 AGAGAACAAAGCCTGATGGTAGAACGAAAGACAGAAATTGCAATAATAAGGATATTCTAGTGGAAACAA
 AAGTATTAATTTCAGAAGATAGTATTAAGAATTCTAGTAAAGCAAGTAGTGTGATCAAGAAGAAGTGGAT
 CGCGATGAATCATCATCTCAAAAGCAAATGATGGAAAAAGGCCACAGTAAGCCTAAAAGGAACCT
 35 TCCTAAAACAGGAGATGCCACTCAGACTGTAAAGCATCTACGGGAGGGATTATTCTGTATCAT
 TAAGTTTTACAATAAGAAAATGAAACTTTAT

SEQ ID NO. 58

MKVKNKILTMVALTVLTCATYSSIGYADTSKNTDTSVVTTLSEEKRSDELDQSSTGSSSENESSSS
 40 SEPETNPSTNPPTTEPSQPSPSEENKPDGRTKTEIGNNKDISSGKVLISEDSIKNFSKASSDQEEDV
RDESSSSKANDGKKGHSKPKKELPKTGDSHSDTVIASTGGIILLSLSFYNKKMKLY

GBS 4 contains an N-terminal leader or signal sequence which is underlined at the beginning of SEQ ID NO: 58 above. In one embodiment, one or more amino acids from the N-terminal leader or signal peptide domain of GBS 4 are removed. An example of such a GBS 4 fragment is set forth below as SEQ ID NO 59.

SEQ ID NO 59

DTSDKNTDTSVVTTLSEEKRSDELDQSSTGSSSENESSSEPETNPSTNPPTTEPSQPSPSEENKP
DGRTKTEIGNNKDISSGKVLISEDSIKNFSKASSDQEEVDRDESSSKANDGKKGHSKPKKELPKTG
DSHSDTVIASTGGIILLSLSFYNKKMKLY

5

A further N-terminal section of GBS 4 may be removed to facilitate recombinant expression.

An example of such a GBS 4 fragment is set forth below as SEQ ID NO: 60.

SEQ ID NO: 60

10 DQSSTGSSSENESSSEPETNPSTNPPTTEPSQPSPSEENKP
DGRTKTEIGNNKDISSGKVLISEDSIKNFSKASSDQEEVDRDESSSKANDGKKGHSKPKKELPKTG
DSHSDTVIASTGGIILLSLSFYNKK
MKLY

15 GBS 4 contains an C-terminal transmembrane region which is underlined at the end of SEQ
ID NO: 58 above. In one embodiment, one or more amino acids from the C-terminal transmembrane
region is removed. An example of such a GBS 4 fragment is set forth below as SEQ ID NO: 61.

SEQ ID NO: 61

20 MKVKNKILTMVALTVLTCATYSSIGYADTS
DKNTDTSVVTTLSEEKRSDELDQSSTGSSSENESSSE
SEPETNPSTNPPTTEPSQPSPSEENKP
DGRTKTEIGNNKDISSGKVLISEDSIKNFSKASSDQEEVDR
DESSSKANDGKKGHSKPKKE

25 In one embodiment, both the N-terminal leader or signal domain and the C-terminal
transmembrane domain are removed from the GBS 4 sequence. An example of such a GBS 4
fragment is set forth below as SEQ ID NO: 62.

SEQ ID NO: 62

30 DTSDKNTDTSVVTTLSEEKRSDELDQSSTGSSSENESSSEPETNPSTNPPTTEPSQPSPSEENKP
DGRTKTEIGNNKDISSGKVLISEDSIKNFSKASSDQEEVDRDESSSKANDGKKGHSKPKKE

In yet another embodiment, the N-terminal leader or signal domain, a further N-terminal
region and the C-terminal transmembrane domain are removed from the GBS 4 sequence. An
example of such a GBS 4 fragment is set forth below as SEQ ID NO: 63.

35 SEQ ID NO: 63

DQSSTGSSSENESSSEPETNPSTNPPTTEPSQPSPSEENKP
DGRTKTEIGNNKDISSGKVLISEDSIKNFSKASSDQEEVDRDESSSKANDGKKGHSKPKKE

GBS 22

40 GBS 22 refers to a putative adhesion lipoprotein. Nucleotide and amino acid sequences of
GBS 22 sequenced from serotype V isolated strain 2603 V/R are set forth in Ref. 2 as SEQ 8583 and
SEQ ID 8584. These sequences are set forth below as SEQ ID NOS 64 and 65:

SEQ ID NO. 64

ATGAAAAGGATACGGAAAAGCCTTATTTGTTCTCGGAGTAGTTACCCCTAATTGCTTATGTGCTTG
 5 TACTAAACAAAGCCAGAAAAAAATGGCTTGTCACTAGTAGCTTTATCCAGTATATTCCATTA
 CAAAAGCAGTTCTGGTGATTGAATGATATTAAAATGATTGATCACAGTCAGGTATTCATGGTTT
 10 GAACCTCATCAAGTGTGCTGCCATTATGATGCTGATCTATTCTTATCATTGCACACACT
 AGAACGTTGGCGAGACGTTGGAACCTAGTTGCATCACTCTAAAGTATCTGAATTGAAGCTTCAA
 AAGGTTATGACTTGGATAAAGTTCATGGCTTAGAAGATGTAGAGGCAGAAAAAGGAGTAGATGAGTCA
 ACCTTGTATGACCCCTCACACTGGAATGACCCGTAAAAGTATCTGAGGAAGCACAACTCATCGCTAC
 15 ACAATTAGCTAAAAGGATCTAAAAACGCTAAGGTTATCAAAAAAATGCTGATCAATTAGTGA
 AGGAATGGCTATTGCAGAGAAGTATAAGCCAAAATTAAAGCTGCAAAGTCTAAATAACTTTGTGACT
 TCACATACAGCATTCTCATACCTAGCTAACGCGATACGGATTGACTCAGTTAGGTATTGCAGGTGTC
 AACCGAGCAAGAACCTAGTGTCTAAAAAATTAGCCGAAATTAGGAGTTGTGAAAACATATAAGGTTA
 AGACTATTGCAAGTTAAGTCCTTARAAGCAGTCCCACAAACAAATAAGATTACTTAGAAAATTGGA
 AACTAATCTTAAGGTACTTGTCAAATCGTTAAATCAATAG

SEQ ID NO. 65

MKRI RKS LI FVLGVVTLICLCACTKQSQQKNGLSVVTSFYPVYSITKAVSGDLNDIKMIRSQSGIHGF
 20 EPSS SDVAAIYDADLFLYHSHTLEAWARRLEPSLHHSKVS VIEASKGMLDKVHGLEDEVAEKGVDES
 TLYD PHTWNDPVKVSEEAQLIATQLAKKDPKNAKVYQKNADQFSKAMAIAEKYKPKFKAASKYFVT
 SHTAFSYLAKRYGLTQLGIAGVSTEQEPSAKLAEIQEFVKTYKVKTIFVEEGVSPKLAQAVASATRV
 KIASLSPLXAVPKNNKDYLENLETNLKVLVKSLNQ

GBS 85

25 GBS 85 refers to a putative cell division protein (DivIB). Nucleotide and amino acid
 sequences of GBS 85 sequenced from serotype V isolated strain 2603 V/R are set forth in Ref. 2 as
 SEQ ID 215 and SEQ ID 216. These sequences are set forth below as SEQ ID NOS 66 and 67:

SEQ ID NO. 66

30 ATGCCTAAGAAGAAATCAGATACCCAGAAAAAGAAGAAGTTGTCTAACGGAAATGGCAAAAGCGTAA
 CCTTGAATTTTAAAAAAACGCAAAGAAGATGAAGAAGAACAAAACGTATTAACGAAAAATTACGCT
 TAGATAAAAGAAGTAAATTAAATATTCTCTGAAGAACCTAAAATACTACTAAAATTAGAAG
 35 CTTCATTTCCAAGATTCAAGACCTAACGATTGAAAAGAAACAGAAAAAGAAAAATAGTCACACAG
 CTTAGCCAAAACTAATCGCATTAGAACCTATATTGTAGTAGCATTCTAGTCATTAGTT
 CGTTTTCTACTAACTCCTTTAGTAAGCAAAAACAATAACAGTTAGTGGAAATCAGCATAACACCT
 GATGATATTGATAGAGAAAACGAATATTCAAAAAACGATTATTCTTTCTTTAATTAAACA
 TAAAGCTATTGAACAACGTTAGCTCAGAAGATGTATGGGAAAAACAGCTCAGATGACTTATCAAT
 40 TTCCAATAAGTTCATATTCAAGTTCAAGAAAATAAGATTATTGCATATGCACATACAAAGCAAGGA
 TATCAACCTGTCTGGAAACTGGAAAAAGGCTGATCCTGAAATAGTCAGAGCTACCAAGCACTT
 CTTAACAAATTAACCTGATAAGGAAGATAGTATTAAAGCTATTAAAGATTAAAGGCTTAGACC
 CTGATTTAATAAGTGAGATTCAAGGTGATAAGTTAGCTGATTCTAAAACGACACCTGACCTCTGCTG
 TTAGATATGCACGATGGAAATAGTATTAGAATACCATTATCTAAATTAAAGAAAGACTCCTTTA
 CAAACAAATTAAGAAGAACCTTAAGGAACCTCTATTGTTGATATGGAAGTGGGAGTTACACAACAA
 45 CAAATACCATTAAGCAATCAACCCCTGTTAACAGCAGAACAGATAACAAAAATAACTCAACTGATAAAACACAA
 ACACAAAATGGTCAGGTTGGAAAATAGTCAGGACAAACAAATAACTCAAATACTAATCAACAAAGG
 ACAACAGATAGCAACAGAGCAGGCACCTAACCCCTCAAATGTTAAT

SEQ ID NO. 67

50 MPKKKSDTPEKEEVVLTEWQKRNLFLKKRKEDEEEQKRINEKLRLDKRSKLNISSEEPQNTTKIKK
 LHPKISRPKIEKKQKKEKIVNSLAKTNRIRTAIFVVAFLVILVSFLLTPFSKQKTITVSGNQHTP
 DDILIEKTNIQKNDYFFSLIFKHKAIEQRLAAEDVWVKTAQMTYQFPNKFHIQVQENKIIAYAHTKQG

YQ PVLETGKKADPVNSSELPKHFLTINLDKEDSIKLLIKDLKALDPDLISEIQVISLADSKTPDLLL
LDMHDGNSIRIPLSKFKERLPFYKQIKKNLKEPSIVDMEVGVYTTNTIESTPVKAEDTKNKSTDKTQ
TQNGQVAENSQGQTNNNSNTNQQGQQIATEQAPNPQNVN

5 **GBS 147**

GBS 147 refers to a putative protease. Nucleotide and amino acid sequences of GBS 147 sequenced from serotype V isolated strain 2603 V/R are set forth in Ref. 2 as SEQ ID 8525 and SEQ ID 8526. These sequences are set forth below as SEQ ID NOS 68 and 69.

10 **SEQ ID NO. 68**

GTGGATAAACATCACTCAAAAAGGCTATTTAAAGTTAACACTTATAACAACTAGTATTTATTAAT
GCATAGCAATCAAGTGAATGCAGAGGAGCAAGAATTAAAAACCAAGAGCAATCACCTGTAATTGCTA
ATGTTGCTCAACAGCCATCGCCATCGTAACTACTAATAACTGTTAAAAACATCTGTAACAGCTGCT
TCTGCTAGTAATACAGCGAAAGAAATGGGTGATACATCTGTAACAGAAGATGAATT
15 ATTAGAAGAGTTATCTAAAACCTGATACGTCTAATTGGGGCTGATCTTGAAGAAGAATATCCCT
CTAAACCAGAGACAACAAATAAGAAAAGCAATGTAGTAACAAATGCTTCAACTGCAATAGCACAG
AAAGTTCCCTCAGCATATGAAGAGGTGAAGGCCAGAAAGCAAGTCATCGCTGCTTGTGATACATC
TAAAATAACAAAATTACAAGCCATAACCCAAAGAGGAAAGGGAAATGTAGTAGCTATTATTGATACTG
20 GCTTGTGATATTAACCATGATATTTCTGTTAGATAGCCAAAAGATGATAAGCACAGCTTAAACT
AAGACAGAATTGAGGAATTAAAGCAAAACATAATACACTTATGGAAATGGGTAACGATAAGAT
TGTGTTGACATAACTACGCCAACAAATACAGAAACGGTGGCTGATATTGCAAGCAGCTATGAAAGATG
GTTATGGTCAGAACAGAATATTCGATGGTACACACGTTGCTGGTATTTGTAGGTAATAGT
AAACGTCCAGCAATCAATGGTCTTCTTGTAGAAGGTGCAGCGCAAATGCTCAAGTCTTAAATGCG
25 TATTCCAGATAAAATTGATTGGACAAATTGGTGAAGCATATGCTAAAGCAATCACAGACGCTGTTA
ATCTAGGAGCAAAACGATTAATATGAGTATTGGAAAAACAGCTGATTCTTAAATTGCTCTCAATGAT
AAAGTTAAATTAGCACTAAATTAGCTCTGAGAACGGCGTTGCAGTTGTTGGCTGCCGGAATGA
AGGCAGATTGGTATGGATTATAGCAAACCAATTATCAACTAATCCTGACTACGGTACGGTTAATAGTC
CAGCTATTCTGAAGATACTTGAGTGGCTAGCTATGAATCACTAAACATATCAGTGAGGTCGTT
30 GAAACAACATTGAAGGTAAAGTTAGTTAAGTGGCGATTGTGACTTCTAAACCTTTGACAAGGTAA
GGCCTACGATGTGGTTATGCCAATTATGGTGCACAAAAAGACTTGAAGGTAAAGGACTTAAAGGTA
AGATTGCATTAATTGAGCGTGGTGGTGGACTTGATTTATGACTAAACACTCATGCTACAAATGCA
GGTGGTTGGTATCGTTATTAAACGATCAAGAAAAACGTGAAATTCTAATTCCCTACCGTGA
ATTACCTGTGGGATTATTAGTAAAGTAGATGGCGAGCGTATAAAACTATCAAGTCAGTTAACAT
TTAACCAAGAGTTGAAGTAGTTGATAGCCAAGGTGGTAATCGTATGCTGGACAATCAAGTTGGGC
35 GTGACAGCTGAAGGAGCAATCAAGCTGATGTAACAGCTTCTGGCTTGAATTATTCTCAACCTA
TAATAATCAATACCAAACAATGTCTGGTACAAGTATGGCTTACCCACATGTCAGGATTAATGACAA
TGCTTCAAAGTCATTGGCTGAGAAATATAAAGGGATGAATTAGATTCTAAAAATTGCTAGAATTG
TCTAAAAACATCCTCATGAGCTCAGCACAGCATTATATAGTGAAGAGGATAAGCGTTTATTCA
ACGTCAGCAAGGTGCAGGTGTTGATGCTGAAAGCTATCCAAGCTCAATATTATATTACTGGAA
40 ACGATGGCAAAGCTAAAATTATCTCAAACGAATGGGAGATAATTGATATCACAGTTACAATT
AAACTTGTAGAAGGTGTCAAAGAATTGTATTATCAAGCTAATGTAGCAACAGAACAGTAATAAAGG
TAAATTGCCCTAAACCACAAGCCTGCTAGATACTAATTGGCAGAAAGTAATTCTCGTGATAAAAG
AAACACAAGTCGATTTACTATTGATGCTAGTCATTTAGTCAGAAATTAAAGAACAGATGGCAAAT
GGTTATTCTTAGAAGGTTTGTACGTTAAAGAAGCCAAGGATAGTAATCAGGAGTTAATGAGTAT
45 TCCTTTGTAGGATTAATGGTATTGCGAACCTACAAGCACTTGAACACCGATTATAAGACGC
TTCTAAAGGTAGTTCTACTATAACCAAATGATACAACCTACATAAGACCAATTGGAGTACAATGAA
TCAGCTCCTTGTAAAGCAACAACTATACTGCCTGTTAACACAATCAGCGTCTGGGCTATGTTGA
TTATGTCAAAATGGTGGGAGTTAGAATTAGCACCAGGAGAGTCCAAAAGAATTATTAGGAACCTT
TTGAGAATAAGGTGAGGATAAAACAATTGATCTTTGGAAAGAGGATGCAGCGAATAATCCATATT
50 GCCATTCTCCAAATAAAGATGGAAATAGGGACGAAATCACTCCCCAGGCAACTTCTTAAGAAATGT
TAAGGATATTCTGCTCAAGTTAGATCAAAATGGAAATGTTATTGGCAAAGTAAGGTTTACCAT

5 CTTATCGTAAAAATTCCATAATAATCCAAAGCAAAGTGTGGTCATTATCGTATGGATGCTCTTCAG
 TGGAGTGGTTAGATAAGGATGGCAAAGTTGTAGCAGATGGTTTATACCTATCGCTTACGTTACAC
 ACCAGTAGCAGAAGGAGCAAATAGTCAGGAGTCAGACTTAAAGTACAAGTAAGTACTAACCAA
 ATCTTCCTTCACGAGCTCAGTTGATGAAACTAATCGAACATTAAGCTAGCCATGCCTAACCAA
 AGTTATGTTCTACATATCGTTACAATTAGTTATCTCATGTTAAAAGATGAAGAATATGGGGA
 10 TGAGACTCTTACCATTTCCATATAGATCAAGAAGGTAAGTGACACTCCTAAACGGTTAAGA
 TAGGAGAGAGTGAGGTTGCGGTAGACCTAACGGCTTGACACTGTTGTGGAAGATAAGCTGGTAAT
 TTCGCAACGGTAAATTGTCGATCTCTGAATAAGGCAGTAGTATCAGAGAAAGAAAACGCTATAGT
 AATTCTAACAGTTCAAATATTTGATAACTGAAAAAGAACCTATGTTATTCCTAAAAAGAAA
 15 10 AAGTAGTAAACAAGAATCTAGAAGAAATAATATTAGTTAAGCCGAAACTACAGTTACTACTCAATCA
 TTGCTAAAGAAATAACTAAATCAGGAAATGAGAAAGTCCTCACTTCTACAAACAATAATAGTAGCAG
 AGTAGCTAAAGATCATATCACCTAACATAACGGGATTCTGTTAACCATACCTACCTAGTACATCAG
 ATAGAGCAACGAATGGCTATTGTTGACTTGGCATTGTTATCTAGTTACTCTTATTTGAAA
 CCCAAAAAGACTAAAAATAATAGTAAA

SEQ ID NO. 69

20 VDKHHSKKAIKLTLITTSILLMHSNQVNAAEQELKNQEQSPVIANVAQQPSPSVTTNTVEKTSVTAAS
 SASNTAKEMGDTSVKNDKTEDELLEELSKNLDTSNLGADLEEEYPSKPETTNKESNVVTNASTAIAQ
 KVPSAYEEVKPESKSSLAVLDTSKITKLQAITQRGKGNVVAIIDTGFIDNHDFRLDSPKDDKHSFKT
 KTEFEELKAKHNITYGKVNNDKIVFAHNYANNTETVADIAAAAMKDGYGSEAKNISHGTHVAGIFVGNS
 KRPAINGLLEGAAPNAQVLLMRIPDKIDSDKFGEAYAKAITDAVNLGAKTINMSIGKTADSLIALND
 KVKLALKLASEKGVAVVVAAGNEGAFGMDYSKPLSTNPDYGTVNSPAISEDTLSVASYESLKTISEVV
 ETTIEGKLVKLPIVTSKPFDKGKAYDVVYANYGAKDDEGKDFKGKIALIERGGGLDFMTKITHATNA
 GVGIVIFNDQEKRGNFLIPYRELPGIISKVDERIKNTSSQLTFNQSFEVVDQSQGGNRMLEQSSWG
 25 VTAEGAIKPDTASGFEIYSSTYNNQYQTMMSGTSMASPHVAGLMTMLQSHLAEKYKGMNLDSSKKLEL
 SKNILMSSATALYSEEDKAFYSPRQQGAGVVDAEKAIQAQYYITGNDGKAKINLKRMDKFDTVTIH
 KLVEGVKELYQANVATEQVNKGKFALKPQALLDTNWQKVILRDKETQVRFTIDASQFSQKLKEQMAN
 GYFLEGFVRFKEAKDSNQELMSIPFGNGDFANLQALETPIYKTLSKGSFYYPNDTTHKDQLEYNE
 SAPFESNNYTALLTQSASWGVYDVVKNGGELELAPESPKRIILGTFENKVEDKTIHLERDAANNPYF
 30 AISPNKDGNRDEITPQATFLRNVKDISAQVLDQNGNVIWQSKVLPSPYRKNFHNNPKQSDGHYRMDALQ
 WSGLDKGKVVADGFYTYRLRYTPVAEGANSQESDFKVQVSTKSPNLPSPRAQFDETNRTLSLAMPKES
 SYVPTYRLQLVLSHVVKDEEYDGETSYHYFIDQEGKVTLPKTVKIGESEVAVDPKALTIVVEDKAGN
 FATVKLSDLNKAJVSEKENAIVISNSFKYFDNLKKEPMFISKKEVVKKNLEEIIILVKPQTTVTTQS
 35 LSKEITKSGNEKVLSTNNNSRVAKISPKHNGDSVNHTLPSTS RATNGLFVGTLLALLSSLLYLK
PKKTKNNSK

GBS 147 contains an N-terminal leader or signal sequence region which is indicated by the underlined sequence at the beginning of SEQ ID NO 69 above. In one embodiment, one or more amino acids from the leader or signal sequence region of GBS 147 are removed. An example of such a GBS 147 fragment is set forth below as SEQ ID NO: 70.

SEQ ID NO: 70

40 EEQELKNQEQSPVIANVAQQPSPSVTTNTVEKTSVTAASASNTAKEMGDTSVKNDKTEDELLEELSKN
 LDTSNLGADLEEEYPSKPETTNKESNVVTNASTAIAQVPSAYEEVKPESKSSLAVLDTSKITKLQA
 ITQRGKGNVVAIIDTGFIDNHDFRLDSPKDDKHSFKTKEFEELKAKHNITYGKVNNDKIVFAHNYA
 NNTEVADI AAAMKDGYGSEAKNISHGTHVAGIFVGNSKRPAINGLLEGAAPNAQVLLMRIPDKIDS
 DKFGEAYAKAITDAVNLGAKTINMSIGKTADSLIALNDVKLALKLASEKGVAVVVAAGNEGAFGMDY
 SKPLSTNPDYGTVNSPAISEDTLSVASYESLKTISEVVETTIEGKLVKLPIVTSKPFDKGKAYDVVYA
 NYGAKKDFEGKDFKGKIALIERGGGLDFMTKITHATNAGVVGIVIFNDQEKRGNFLIPYRELPGIIS
 50 KVDGERIKNTSSQLTFNQSFEVVDQSQGGNRMLEQSSWGVTAEAGAIKPDTASGFEIYSSTYNNQYQTM
 SGTSMASPHVAGLMTMLQSHLAEKYKGMNLDSSKKLELSKNILMSSATALYSEEDKAFYSPRQQGAGV

5 VDAEKAIQAQYYITGNDGAKINLKRMDKFIDTVTIHKLVEGVKELYQANVATEQVNKGKFALKPQ
 ALLDTNWQKVILRDKETQVRFTIDASQFSQKLKEQMANGYFLEGFVRFKEAKDSNQELMSIPFVGFNG
 DFANLQALETPIYKTL SKGSFYYKPNDTTHKDQLEYNESAPFESNNYTALLTQSASWGYVDYVKNGGE
 LELAPE SPKRIILGT FENKVEDKTIHLLE RDAANNPYFAISPNKDGRNDEITPQATFLRNVKDISAQV
 LDQNGNVIWQSKVLP SYRKNFHNNPKQSDGHYRMDALQWSGLDKDGKVVA DGFYTYRLRYTPVAEGAN
 10 SQESDFKVQVSTKS PNLP SRAQFDETNRTLSLAMPKESYYVPTYRLQLVL SHVVKDEEYGD ETSYHYF
 HIDQEGKVTL PKTVKIGESEVAVDPKALT LVVEDKAGNFATVKLSDLLNKAVVSEKENAIVISNSFKY
 FDNLKKEPMFISKKEKVNKNLEEIIILVKPQTTQSLSK EITKSGNEKVL TSTNNNSSRVAKIIISP
 KHNGDSVNHTLPSTS DRATNGLFVGTLALLSSLLYLKP KKT KNN SK

10 **GBS 147** also contains a C-terminal transmembrane and/or cytoplasmic region which may be located within the underlined sequence near the end of SEQ ID NO: 69 above. In one embodiment, one or more amino acids from the transmembrane and/or cytoplasmic region are removed. An example of such a GBS 147 fragment is set forth below as SEQ ID NO: 71.

15 **SEQ ID NO: 71**
 VDKHHS KKAILKLT LITTS ILLMHSNQVNAEEQELKNQE QSPVIANVAQQPSPSVTTNTVEKTSVTAA
 SASNTAKEMGDT SVKNDKTE DELLEELSKN LDT S NLGADLEEEYPSK PETTNNKESNVVTNASTAIAQ
 20 KVPSAYEEVKPESKSSLAVL DTSK ITKLQAI TQRGKGNVVAI IDTGF D INHDI FRLD SPKDDKHS FKT
 KTE FEELKAKHNITYGK WNDKIVFAHNYANNTETVADIAAA M KDG YGSEAKNISHGTHVAGI FVGNS
 KRPAI NGLLEGAAPNAQVLLM RIPDKIDS DKFGEAYAKAI TDAVNLGAKTINMSIGKTADSLIALND
 KVKLALKLASEKG VAVVVAAGNEAGFMDY SKPLS TNP D YGT VNNSPAI E DTL SVAS YES LKT ISEVV
 25 ETTIEGKLV KLP I VTSK PFDKG KAYD VVY ANYGAKKDFEGKDFKG KIALIERGGGLDFMTK ITHATNA
 GVGIVI FNDQEK RGNFLI PYREL P VGI IS KV DGERIK NTSS QLT FNQS FEV VD S QGGN RMLE QSSWG
 VTAEGA I KPDVTAS GFEI YSSTYNNQYQTM GTS MAS PHVAGL MTL QSHLAE KYKG MNL DSK KLEL
 30 SKN ILMSSA TALYSEEDKA FYS PRQQGAGV DAEKAI QAQYYITGNDGAKINLKRMDKFIDTVTIH
 KLV EGV KELYQANVATEQVNKGKFALKPQALLDTNWQKVILRDKETQVRFTIDASQFSQKLKEQMAN
 GYFLEG FVRFKEAKDSNQELMSIPFVGFNGDFANLQALETPIYKTL SKGSFYYKPNDTTHKDQLEYNE
 35 SAPFESNNYTALLTQSASWGYVDYVKNGGELELAPE SPKRIILGT FENKVEDKTIHLLE RDAANNPYF
 AISPNKDG NRDEITPQATFLRNVKDISAQVLDQNGNVIWQSKVLP SYRKNFHNNPKQSDGHYRMDALQ
 WSGLDKDGKVVA DGFYTYRLRYTPVAEGANSQESDFKVQVSTKSPNLP SRAQFDETNRTLSLAMPKES
 SYVPTYRLQLVL SHVVKDEEYGD ETSYHYF HIDQEGKVTL PKTVKIGESEVAVDPKALT LVVEDKAGN
 FATVKLSDLLNKAVVSEKENAIVISNSFKYFDNLKKEPMFISKKEKVNKNLEEIIILVKPQTTQSL
 LSKEITKSGNEKVL TSTNNNSSRVAKIIISP KHN GDSVNHT

35 In one embodiment, one or more amino acids from the leader or signal sequence region and one or more amino acids from the transmembrane or cytoplasmic region are removed from the GBS 147 sequence. An example of such a GBS 147 fragment is set forth below as SEQ ID NO 72.

40 **SEQ ID NO: 72**
 EEQELKNQE QSPVIANVAQQPSPSVTTNTVEKTSVTAA SASNTAKEMGDT SVKNDKTE DELLEELSKN
 LDT S NLGADLEEEYPSK PETTNNKESNVVTNASTAIAQKVPSAYEEVKPESKSSLAVL DTSK ITKLQAI
 45 ITQRGKGNVVAI IDTGF D INHDI FRLD SPKDDKHS FKT KTE FEELKAKHNITYGK WNDKIVFAHNYA
 NNNTETVADIAAA M KDG YGSEAKNISHGTHVAGI FVGNSK RPAI NGLLEGAAPNAQVLLM RIPDKIDS
 DKFGEAYAKAI TDAVNLGAKTINMSIGKTADSLIALNDKV K LALKLASEKG VAVVVAAGNEAGFMDY
 SKPLS TNP D YGT VNNSPAI E DTL SVAS YES LKT ISEVVETTIEGKLV KLP I VTSK PFDKG KAYD VVY
 NYGAKKDFEGKDFKG KIALIERGGGLDFMTK ITHATNAGVGIVI FNDQEK RGNFLI PYREL P VGI IS
 KV DGERIK NTSS QLT FNQS FEV VD S QGGN RMLE QSSWG VTAEGA I KPDVTAS GFEI YSSTYNNQYQTM
 SGTSMAS PHVAGL MTL QSHLAE KYKG MNL DSK KLEL SKN ILMSSA TALYSEEDKA FYS PRQQGAGV

VDAEKAI QAQYYITGNDGKAKINLKRMDKFDTVTIHKLVEGVKELYQANVATEQVNKGKFALKPQ
 ALLDTNWQKVILRDKETQVRFTIDASQFSQKLKEQMANGYFLEGFVRFKEAKDSNQELMSIPFVGFNG
 DFANLQALETPYKTL SKGSFYYKPNDTTHKDQLEYNESAPFESNNYTALLTQSASWGYVDYVKNGGE
 LELAPES PKRIILGT FENKVEDKTIHLLE RDAANNPYFAISPNKDGNRDEITPQATFLRNVKDISAQV
 5 LDQNGNVIWQSKVLP SYRKNFHNPKQSDGHYRMDALQWSGLDKDGKVVADGFYTYRLRYTPVAEGAN
 SQESDFKVQVSTKSPNLP SRAQFDETNR TL SLAMPKESSYVPTYRLQLVLSHVVKDEEYGDTSYHYF
 HIDQEGKVTLPKTVKIGESEVA VDPKALT LVEDKAGNFATVKLSDLLNKA VVSEKENAIVISNSFKY
 FDNLKKE PMFISKKEV VVNKNLEE IILVKPQTTQSL SKEITKSGNEKVLTSTNNN S R VAKI I SP
 KHNGDSVNHT

10

GBS 173

GBS 173 refers to an amidase family protein. Nucleotide and amino acid sequences of GBS
 173 sequenced from serotype V isolated strain 2603 V/R are set forth in Ref. 2 as SEQ ID 8787 and
 SEQ ID 8788. These sequences are set forth below as SEQ ID NOS 73 and 74:

15

SEQ ID NO. 73

ATGAAACGTAAATACTTATTCTTAATACGGTACGGTTAACGTTAGCTGCTGCAATGAATACTAG
 CAGTATCTATGCTAATAGTACTGAGACAAGTGCTTCAGTAGTTCCTACTACAAATACTATCGTCAAA
 20 CTAATGACAGTAATCCTACCGCAAAATTGTATCAGAACATCAGGACAATCTGTAATAGGTCAAGTAAAA
 CCAGATAATTCTCGGGCGCTTACAACAGTTGACACGCCCTCATCATATTCAGCTCCAGATGCTTAAA
 AACAACTCAATCAAGTCTGCTGTTGAGAGTACTTCTACTAAGTTACTGAAGAGACTTACAAACAAA
 AAGATGGTCAAGATTAGCCAACATGGTGAGAAGTGGTCAAGTTACTAGTGAGGAACCTGTTAATATG
 GCATACGATATTATTGCTAAAGAAAACCCATCTTAAATGCAGTCATTACTACTAGACGCCAAGAAC
 25 TATTGAAGAGGCTAGAAA ACTTAAAGATAACCAATCAGCCGTTTTAGGTGTTCCCTGTTAGTCAAGG
 GGTTAGGGCACAGTATTAAAGGTGGTGAACCAATAATGGCTTGATCTATGCAGATGGAAAAATTAGC
 ACATTGACAGTAGCTATGTCAAAAAATATAAAGATTAGGATTATTATTAGGACAAACGAACCT
 TCCAGAGTATGGGTGGCGTAATATAACAGATTCTAAATTACGGTCTAACGCATAATCCTGGGATC
 TTGCTCATATGCTGGTGGCTCTCTGGTGGAAAGTGCAGCAGCCATTGCTAGCGGAATGACGCCAATT
 GCTAGCGGTAGTGATGCTGGTGGTTCTATCCGTATTCCATCTTCTGGACGGGCTGGTAGGTTAAA
 30 ACCAACAAAGAGGATTGGTGAGTAATGAAAAGCCAGATTGGTATAGTACAGCAGTTCAATTCCATTAA
 CTAAGTCATCTAGAGACGCAGAACATTATTAACCTTATCTAAAGAAAAGCGATCAAACGCTAGTATCA
 GTTAATGATTAAATCTTACCAATTGCTTACTTTGAAATCCAATGGGAACAGAAGTTAGTCA
 AGATGCTAAAACGCTATTATGGACAACGTCAACATTCTTAAGAAAACAAGGATTCAAAGTAACAGAGA
 TAGACTTACCAATTGATGGTAGAGCATTATGCGTGATTATTCAACCTGGCTATTGGCATGGGAGGA
 35 GCTTTTCAACAATTGAAAAAGACTAAAAACATGGTTTACTAAAGAAGACGTTGATCCTATTAC
 TTGGCAGTTCATGTTATTATCAAATTCAAGATAAGGCTGAACCTAAGAAATCTATTATGGAAGCCC
 AAAAACATATGGATGATTATCGTAAGGCAATGGAGAAGCCTCACAAGCAATTCCCTATTCTTATCG
 CCAACGACCGCAAGTTAGCCCTCTAAATACAGATCCATATGTAACAGAGGAAGATAAAAGACGAT
 TTATAATATGGAAA ACTTGAGCCAAGAAGAAAATTGCTCTTTAATGCCAGTGGGAGCCTATGT
 40 TGCCTAGAACACCTTTACACAAATTGCTAATATGACAGGACTCCAGCTATCAGTATCCGACTTAC
 TTATCTGAGTCTGGTTACCCATAGGGACGATGTTAATGGCAGGTGCAAACATATGATATGGTATTAA
 TAAATTGCAACTTCTTTGAAAAACATCATGGTTTAAATGTTAAATGGCAAGAATAATAGATAAAG
 AAGTGAAGAAAATTCAACAGTTACTCAAGTATCTATCTAAAGCTCATTCACTTAGTA
 45 AATTAGAAGAAAATTCAACAGTTACTCAAGTATCTATCTAAAGCTCATTCACTTAGTA
 AAATAAACCATCCGTAATGGCATATCAAAAGCACTCCAAAACAGGTGATACAGAATCAAGCCTAT
 CTCCAGTTTAGTAGTAACCCCTTTATTAGCTTGTAGCTTTGTAACAAAAAGAATCAGAAAAGT

SEQ ID NO. 74

5 MKRKYFILNTVTVLTLAAAMNTSS IYANSTETSASVVPTTNTIVQTNDNSNPTAKFVSESGQSVIGQVK
 PDNSAALT TVDTPHHISAPDALKTTQSSPVVESTSTKLTEETYKQKDGQDLANMVRSGQVTSEELVN
 AYDIIAKENPSLNAVITTRRQEAIIEEARKLKDTNQPFLGVPLLVKG~~L~~HSIKGGETNNGLIYADGKIS
 10 TFDSSYVKKYKDLGFIILGQTNFPEYGWRNITDSKLYGLTHNPWDLAHNAGGSSGGSAAAIASG~~M~~TP
 ASGSDAGGSIRIPSSWTGLVGLKPTRGLVSNEKPD~~S~~YSTAVHFPLTKSSRDAETLLTYLKKSDQTLVS
 VNDL~~K~~SLPIAYTLKSPMGTEVSQDAKNAIMDNVTFLRKQGFKVTEIDLPIDGRALMRD~~Y~~STLAIGMGG
 AFSTIEKDLKKHGFTKEDVD~~P~~ITWAHVVIYQNSDKAELKKSIMEAQKHMDDYR~~K~~AMEKLHKQFPIFLS
 15 PTTASLAPLNTDPYVTEEDKRAIYNMENLSQEE~~R~~IALFNRQWEPM~~L~~RRTPFTQIANMTGLPAISIPTY
 LSESGLPIGTMLMAGANYDMVLIK~~F~~FEKHGFNVK~~W~~QRIIDKEV~~K~~PSTGLIQPTNSL~~F~~KAHSSLV
 NLEENSQVTQVSISKKWMKSSVKNKPSVMAYQ~~K~~ALPKTG~~D~~TESSLS~~P~~VLVV~~T~~LLACFS~~F~~VTKKNQKS

GBS 173 contains an N-terminal leader or signal sequence region which is indicated by the

15 underlined sequences at the beginning of SEQ ID NO: 74 above. In one embodiment, one or more
 amino acids from the leader or signal sequence of GBS 173 are removed. An example of such a GBS
 173 fragment is set forth below as SEQ ID NO: 75.

SEQ ID NO: 75

20 TTNTIVQTNDNSNPTAKFVSESGQSVIGQVKPDNSAALT TVDTPHHISAPDALKTTQSSPVVESTSTKL
 TEETYKQKDGQDLANMVRSGQVTSEELVN~~MAYDI~~IAKENPSLNAVITTRRQEAIIEEARKLKDTNQPFL
 GVPLLVKG~~L~~HSIKGGETNNGLIYADGKISTFDSSYVKKYKDLGFIILGQTNFPEYGWRNITDSKLYG
 LTHNPWDLA~~H~~HNAGGSSGGSAAAIASG~~M~~TPIASGSDAGGSIRIPSSWTGLVGLKPTRGLVSNEKPD~~S~~
 TAVHFPLTKSSRDAETLLTYLKKSDQTLVSVNDL~~K~~SLPIAYTLKSPMGTEVSQDAKNAIMDNVTFLRK
 25 QGFKVTEIDLPIDGRALMRD~~Y~~STLAIGMGGAFSTIEKDLKKHGFTKEDVD~~P~~ITWAHVVIYQNSDKAEL
 KKSIMEAQKHMDDYR~~K~~AMEKLHKQFPIFLSPTTASLAPLNTDPYVTEEDKRAIYNMENLSQEE~~R~~IALF
 NRQWEPM~~L~~RRTPFTQIANMTGLPAISIPTYLSESGLPIGTMLMAGANYDMVLIK~~F~~FEKHGFNVK
 WQRIIDKEV~~K~~PSTGLIQPTNSL~~F~~KAHSSLV~~N~~LEENSQVTQVSISKKWMKSSVKNKPSVMAYQ~~K~~ALPKT
 30 GDTESSLS~~P~~VLVV~~T~~LLACFS~~F~~VTKKNQKS

GBS 173 may also contain a C-terminal transmembrane and/or cytoplasmic region which
 may be located within the underlined region near the end of SEQ ID NO: 74 above. In one
 embodiment, one or more amino acids from the transmembrane or cytoplasmic region of GBS 173 are
 removed. An example of such a GBS 173 fragment is set forth below as SEQ ID NO: 76.

35

SEQ ID NO: 76

40 MKRKYFILNTVTVLTLAAAMNTSS IYANSTETSASVVPTTNTIVQTNDNSNPTAKFVSESGQSVIGQVK
 PDNSAALT TVDTPHHISAPDALKTTQSSPVVESTSTKLTEETYKQKDGQDLANMVRSGQVTSEELVN
 AYDIIAKENPSLNAVITTRRQEAIIEEARKLKDTNQPFLGVPLLVKG~~L~~HSIKGGETNNGLIYADGKIS
 45 TFDSSYVKKYKDLGFIILGQTNFPEYGWRNITDSKLYGLTHNPWDLAHNAGGSSGGSAAAIASG~~M~~TP
 ASGSDAGGSIRIPSSWTGLVGLKPTRGLVSNEKPD~~S~~YSTAVHFPLTKSSRDAETLLTYLKKSDQTLVS
 VNDL~~K~~SLPIAYTLKSPMGTEVSQDAKNAIMDNVTFLRKQGFKVTEIDLPIDGRALMRD~~Y~~STLAIGMGG
 AFSTIEKDLKKHGFTKEDVD~~P~~ITWAHVVIYQNSDKAELKKSIMEAQKHMDDYR~~K~~AMEKLHKQFPIFLS
 PTTASLAPLNTDPYVTEEDKRAIYNMENLSQEE~~R~~IALFNRQWEPM~~L~~RRTPFTQIANMTGLPAISIPTY
 45 LSESGLPIGTMLMAGANYDMVLIK~~F~~FEKHGFNVK~~W~~QRIIDKEV~~K~~PSTGLIQPTNSL~~F~~KAHSSLV
 NLEENSQVTQVSISKKWMKSSVKNK

In one embodiment, one or more amino acids from the leader or signal sequence region and one or more amino acids from the transmembrane or cytoplasmic region are removed. An example of such a GBS 173 fragment is set forth below as SEQ ID NO: 77.

5 **SEQ ID NO: 77**

TTNTIVQTNDSNPTAKFVSESGQSVIGQVKPDNSAALTVDTPHISAPDALKTTQSSPVVESTSTKL
 TEETYKQKDGQDLANMVRSGQVTSEELVN MAYDIIAKENPSLNAVITRRQEAI EARKLKDTNQPFL
 GVP LLVKG LGH SIK GGETN NGLIYADG KISTFDSSYVKKYKDLGFIILGQTNFPEYGRNITDSKLYG
 10 LTHNPWDLAHNAGGSSGGSAAAIASG MTPIASGSDAGGSIRIPSSWTGLVGLKPTRGLVSNEKPD SYS
 TAVHFPLTKSSRDAETLLTYLKKSDQTLVSVNDLKS LPIAYTLKSPMGTEVSDAKNAIMDNVTFLRK
 QGF KVTEIDLPI DGR ALMRD YSTLAIGMGGAFSTIEKDLKKHGFTKEDVDPI TWAVHVIYQNSDKAEL
 KKSIMEAQKHMDDYRKAMEKLHKQFPIFLSPTTASLAPLNTDPYVTEEDKRAIYNMENLSQEE RIALF
 NRQWE PMLRRT PFTQIANMTGLPAISIPTYLSESGLPIG TMLMAGANYDMVL IKFAT FFEKHHGFNVK
 15 WQRIIDKEVKPSTGLI QPTNSLFAHSSLVNEEN SQVTQVSISKWMKSSVKNK

15 **GBS 313**

Nucleotide and amino acid sequences of GBS 313 sequenced from serotype V isolated strain 2603 V/R are set forth in Ref. 2 as SEQ ID 4089 and SEQ ID 4090. These sequences are set forth as SEQ ID NOS 78 and 79 below:

20

SEQ ID NO. 78

ATGAAACGTATTGCTGTTAACTAGTGGTGGTGACGCCCTGGTATGAACGCTGCTATCCGTGCAGT
 TGTCGTAAGCAATTCTGAAGGTATGGAAGTTACGGCATCAACCAAGGTTACTATGGTATGGTGA
 25 CAGGGGATATTTCCCTTGGATGCTAATTCTGGTGGGGATACTATCAACCGTGGAGGAACGTTTTA
 CGTT CAGCACGTTATCCTGAATTGCTGAACCTGAAGGT CAGCTAAAGGGATTGAACAGCTAAAAA
 ACACGGTATTGAAGGTGAGTAGTTATCGGTGGTGATGGTTCTTATCATGGT GCTATCGCTTA ACTG
 AGCACGGTTCC CAGCTGTTGGTTGCCGGTACAATTGATAACGATATCGTGGCACTGACTATACT
 ATTGGTTTGACACAGCAGTTGCGACAGCAGTTGAGAATCTTGACCGTCTCGT GATA CATCAGCAAG
 30 TCATAACCGTACTTTGTTGAGGTATGGGAGAAATGCAGGAGATATCGCTCTTGGTCAGGTA
 TCGCTGCAGGTGCAGATCAAATTATTGTCCTGAAGAAGAGTTCAATATTGATGAAGTTGCTCAAAT
 GTTAGAGCTGGCTATGCAGCTGGTAAACATCACCAATCATCGTCCTGCGAGAAGGTGTTATGAGTGG
 TGATGAGTTGCAAAAACAATGAAAGCAGCAGGAGACGATAGCGATCTCGTGTGACGAATTAGGAC
 35 ATCTGCTCCGTGGTAGTCGACGGCTCGT GATCGTCTTAGCATCTCGTATGGGAGCGTACGCT
 GTTCAATTGTTGAAAGAAGGTCGTGGTGGTTAGCCGTTGGTGTCCACAACGAAGAAATGGT GAAAG
 TCCAATT TAGGTTAGCAGAAGAAGGTGCTTGTGACTGATGAAGGAAAATCGTTGTTA
 ATAATCCGCATAAAGCGGACCTTCGCTGGCAGCACTTAATCGTGACCTGCCAACCAAAGTAGTAAA

SEQ ID NO. 79

MKRIAVL TS GGDAPGM NAI RAVVRKAISEGMEVYGINQGYYGMVTGDI FPLDANSVGDTI NRGGTFL
 40 RSARYPEFAELEGQLKGIEQLKKHGIEGVV VIGGDGSYHGAMRLTEHGFPAVGLPGTIDNDIVGTDYT
 IGFDTAVATAVENLDRRLRTSASHNRTFVVEVMGRNAGDIALWSGIAAGADQIIVPEEEFNIDEVVS N
 VRAGYAAGKHHQIIVLAEGVMSGDEFAKTMKAAGDDSDLRVTNLGHLLRGGSPTARDRVLASRMGAYA
 VQLLKEGRGGLAVGVHNEEMVESPILGLAEEGALFSLTDEGKIVVNNPHKADLRLAALNRDLANQSSK

GBS 328

GBS 328 belongs to the 5'-nucleotidase family. Nucleotide and amino acid sequences of GBS 328 sequenced from serotype V isolated strain 2603 V/R are set forth in Ref. 2 as SEQ ID 6015 and SEQ ID 6016. These sequences are set forth below as SEQ ID NOS 80 and 81:

SEQ ID NO. 80

ATGAAAAAGAAAATTATTTGAAAAGTAGTGTCTGGTTAGTCGCTGGGACTTCTATTATGTTCTC
 10 AAGCGTGGTCGCGGACCAAGTCGGTGTCCAAGTTAGGCGTCATGGTGCACCTGACA
 ATACTGGAACAGCAAATATGCCTGATGGAAAAGTTGCTAATGCTGGTACTGCTGCTCAATTAGATGCT
 TATATGGATGACGCTCAAAAAGATTCAAACAAACTAACCCCTAATGGTGAAGCATTAGGGTTCAAGC
 AGGCGATATGGTGGAGCAAGTCCAGCCAACCTGGGCTTCTCAAGATGAACCAACTGTCAAAAATT
 15 TTAATGCAATGAATGTTGAGTATGGCACATTGGTAACCATGAATTGATGAAGGGTTGGCAGAATAT
 AATCGTATCGTTACTGGTAAAGCCCCCTGCTCCAGATTCTAATATTAAATAATTACGAAATCATACCC
 ACATGAAGCTGCAAAACAAGAAATTGTAGTGGCAAATGTTATTGATAAAAGTTAACAAACAATTCTT
 ACAATTGGAAGCCTACGCTATTAAAATATTCTGTAAATAACAAAAGTGTGAACGTTGGCTTATC
 20 GGGATTGTCACCAAAGACATCCAAACCTTGTCTACGTTAAAGGAAATTACAAGCTAAAATGTCAAAGCTATTGTTAGTTC
 TGAAGCTGAAACAATCGTTAAATACGCCAAAGAATTACAAGCTAAAATGTCAAAGCTATTGTTAGTTC
 TCGCACATGTACCTGCAACAAGTAAAATGATATTGCTGAAGGTGAAGCAGCAGAAATGATGAAAAAA
 25 GTCAATCAACTCTTCCCTGAAAATAGCGTAGATATTGTCCTTGCTGGACACAAATCATCAATATAACAAA
 TGGTCTTGTGGTAAAACCTGTTAGTACAAGCGCTCTCAAGGAAAAGCCTATGCTGATGTCAGTG
 GTGCTTAGATACTGATAACACAAGATTCTATTGAGACCCCTTCAGCTAAAGTAATTGCAAGTTGCTCCT
 GGTAAAAAAACAGGTAGTGCCGATATTCAAGCCATTGTTGACCAAGCTAATACTATCGTTAAACAAGT
 AACAGAAGCTAAAATGGTACTGCCGAGGTAAAGTGTATGATTACGCGTTCTGTTGATCAAGATAATG
 30 TTAGTCCGGTAGGCAGCCTCATCACAGAGGCTCAACTAGCAATTGCTCGAAAAGCTGCCAGATATC
 GATTTGCCATGACAAATAATGGTGGCATTCTGTGCTGACTTACTCATCAAACCAGATGGAACAATCAC
 CTGGGGAGCTGCACAAGCAGTCAACCTTTGGTAATATCTTACAAGTGTGCAAATTACTGGTAGAG
 ATCTTATAAAGCACTCAACGAACAATACGCCAAAACAAAATTCTCCTCAAATAGCTGGCTG
 CGATACACTACACAGATAATAAAGAGGGCGGGGAAGAAACACCATTAAAGTTGAAAGCTTATAA
 35 ATCAAATGGTGAGGAAATCAATCCTGATGCAAATCAAATTAGTTATCAATGACTTTTATTGGTG
 GTGGTGATGGCTTGCAAGCTCAGAAATGCCAAACTCTAGGAGCCATTAAACCCGATACAGAGGTA
 TTTATGGCCTATATCACTGATTAGAAAAGCTGGAAAAAGTGGCTCCAAATAATAAACCTAA
 AATCTATGTCACTATGAAGATGGTAATGAAACTATTACACAAAATGATGGTACACATAGCATTATTA
 AGAAAACTTATTAGATCGACAAGGAATTGTAGCACAAGAGATTGATCAGACACTTAAACCAA
 40 ACAAAATCAAATCTACAAAATCAACCTGTAACTACAATTCAACAAAACAATTACACCAATTAC
 AGCTATTAAACCCATGAGAAATTATGCCAAACCATCAAACCTCAACTACTGTAAAATCAAACAAATTAC
 CAAAACAAACTCTGAATATGGACAATCATCCTTATGTCGTCTTGGTGTGGACTTATAGGAATT
 GCTTAAATACAAAGAAAAACATATGAAA

SEQ ID NO. 81

MKKIILKSSVLGLVAGTSIMFSSVFAQVGVQVIGVNDFHGALDNTGTANMPDGKVNAGTAAQLDA
 YMDDAQKDFKQTNPNGESIRVQAGDMVGASPANSLLQDEPTVKNFNAMNVEYGTILGNHEFDEGLAEY
 NRIVTGKAPAPDSNINNITKSYPHEAAKQEIVVANVIDKVNKQIPYNWKPYAIKNIPVNNKSVNVGFI
 GIVTKDIPLVLRKNYEQYEFLDEAETIVKYAKELQAKNVKAIVVLAHPATSKNDIAEGEAAEMMKK
 45 VNQLFPENSVDIVFAGHNHQYTNGLGVGKTRIVQALSQGKAYADVRGVLDTDQDFIETPSAKVIAVAP
 GKKTGSADIQAIVDQANTIVKQVTEAKIGTAEVSMITRSDQDNSPVGSILTEAQLAIARKSWPDI
 DFAMTNNGGIRADLLIKPDGTITWGAAQAVQPFGNILQVVEITGRDLYKALNEQYDQKQNFFLQIAGL
 RYTDTNKEGGEETPFKVVKAYKSNGEEINPDAKYKLVINDFLFGGGDGFAFRNAKLLGAINPDTEV
 FMAYITDLEKAGKKVSVPPNNKPKIYVTMKMVNETITQNDGTHSIKKLYLDRQGNIVAQEIVSDLNQ
 50 TKSKSTKINPVTTIHKKQLHQFTAINPMRNYGKPSNSTVKSQLPKTNSEYQOSFLMSVFGVGLIGI
 ALNTKKKHMK

GBS 328 may contain an N-terminal leader or signal sequence region which is indicated by the underlined sequence at the beginning of SEQ ID NO: 81 above. In one embodiment, one or more amino acids from the leader or signal sequence region of GBS 328 are removed. An example of such a GBS 328 fragment is set forth below as SEQ ID NO: 82.

SEQ ID NO: 82

10 HGALDNTGTANMPDGKVANAGTAAQLDAYMDDAQKDFQTNPNGESIRVQAGDMVGASPANSGLLQDE
 PTVKNFNAMNVEYGTGLNHEFDEGLAEYNRIVTGPAPAPDSNINNITKSYPHEAAKQEIVVANVIDKV
 NKQI PYNWKPYAIKNI PVNNKSVNVGFIGIVTKDIPNLVLRKNYEQYEFLEAETIVKYAKELQAKNV
 KAI VVLAHVPATSKNDIAEGEAAEMMKVNQLFPENSVDIVFAGHNHQYTNGLVGKTRIVQALSQGKA
 YADVRGVLDTDTQDFIETPSAKVIAVAPGKKTGSADIQAIVDQANTIVKQVTEAKIGTAEVSVMITRS
 VDQDNVSPVGSLITEAQLAIARKSWPDI DFAMTNNGGIRADLLIKPDGTITWGAAQAVQPGNILQVV
 EITGRDLYKALNEQYDQKQNFFLQIAGLRYTYTDNKEGGEETPFKVVKAYKSNGEEINPDAKYKLVIN
 15 DFL FGGGDGFASFRNAKLLGAINPDTEVF MAYITDLEKAGKKVSPNNKPKIYVTMKMVNETITQNDG
 THS I IKKLYLDRQGNIVAQEIVSDTLNQTKSKTKINPVTTIHKKQLHQFTAINPMRNYGKPSNSTTV
 KSKQLPKTNSEYQSFMSVFGVGLIGALNTKKHMK

GBS 328 may also contain a transmembrane and/or cytoplasmic domain region. In one embodiment, one or more amino acids from the transmembrane and/or cytoplasmic domain region of GBS 328 are removed. An example of such a GBS 328 fragment is set forth below as SEQ ID NO: 83.

SEQ ID NO: 83

25 MKKKIILKSSVLGLVAGTSIMFSSVFADQVGVQVIGVNDEHGALDNTGTANMPDGKVANAGTAAQLDAY
 YMDDAQKDFQTNPNGESIRVQAGDMVGASPANSGLLQDEPTVKNFNAMNVEYGTGLNHEFDEGLAEY
 NRIVTGPAPAPDSNINNITKSYPHEAAKQEIVVANVIDKVNQKIPYNWKPYAIKNI PVNNKSVNVGFI
 GIVTKDIPNLVLRKNYEQYEFLEAETIVKYAKELQAKNVKAI VVLAHVPATSKNDIAEGEAAEMMK
 VNQLFPENSVDIVFAGHNHQYTNGLVGKTRIVQALSQGKAYADVRGVLDTDTQDFIETPSAKVIAVAP
 30 GKKTGSADIQAIVDQANTIVKQVTEAKIGTAEVSVMITRSVDQDNVSPVGSLITEAQLAIARKSWPDI
 DFAMTNNGGIRADLLIKPDGTITWGAAQAVQPGNILQVV EITGRDLYKALNEQYDQKQNFFLQIAGL
 RYTYTDNKEGGEETPFKVVKAYKSNGEEINPDAKYKLVINDFLFGGGDGFASFRNAKLLGAINPDTEV
 FMAYITDLEKAGKKVSPNNKPKIYVTMKMVNETITQNDGTHS I IKKLYLDRQGNIVAQEIVSDTLNQ
 35 TKS SKTKINPVTTIHKKQLHQFTAINPMRNYGKPSNSTTVKS

In one embodiment, one or more amino acids from the leader or signal sequence region and one or more amino acids from the transmembrane or cytoplasmic region of GBS 328 are removed. An example of such a GBS 328 fragment is set forth below as SEQ ID NO: 84.

SEQ ID NO: 84

40 HGALDNTGTANMPDGKVANAGTAAQLDAYMDDAQKDFQTNPNGESIRVQAGDMVGASPANSGLLQDE
 PTVKNFNAMNVEYGTGLNHEFDEGLAEYNRIVTGPAPAPDSNINNITKSYPHEAAKQEIVVANVIDKV
 NKQI PYNWKPYAIKNI PVNNKSVNVGFIGIVTKDIPNLVLRKNYEQYEFLEAETIVKYAKELQAKNV
 KAI VVLAHVPATSKNDIAEGEAAEMMKVNQLFPENSVDIVFAGHNHQYTNGLVGKTRIVQALSQGKA
 YADVRGVLDTDTQDFIETPSAKVIAVAPGKKTGSADIQAIVDQANTIVKQVTEAKIGTAEVSVMITRS
 45 VDQDNVSPVGSLITEAQLAIARKSWPDI DFAMTNNGGIRADLLIKPDGTITWGAAQAVQPGNILQVV

5 EI TGRDLYKALNEQYDQKQNFLOIAGLRYTYTDNKEGGEETPFKVVKAYKSNSEEINPDALKLVIN
 DFLFGGGDGFAFRNAKLLGAINPDTEVFMAITDLEKAGKKVSPNNPKIYVTMNMVNETITQNDG
 THSIIKKLYLDRQGNIVAQEIVSDTLNQTKSKSTKINPVTTIHKKQLHQFTAINPMRNYGKPSNSTTV
 KS

10 **GBS 656**

GBS 656 refers to a putative DNA-entry nuclease. Nucleotide and amino acid sequences of GBS 656 sequenced from serotype V isolated strain 2603 V/R are set forth in Ref. 2 as SEQ ID 9323 and SEQ ID 9324. These sequences are set forth below as SEQ ID NOS 85 and 86:

15 **SEQ ID NO. 85**

ATGAAAAGATTACATAAACTGTTATAACCGTAATTGCTACATTAGGTATGTTGGGGTAATGACCTT
 TGGTCTTCCAACGCAGCCGAAAACGTAACGCCGATAGTACATGCTGATGTCAATTCACTGTGATA
 CGAGCCAGGAATTCAAAATAATTAAAAATGCTATTGGTAACCTACCATTCAATATGTTAATGGT
 ATTATGAATTAAATAATCAGACAAATTAAATGCTGATGTCAATGTTAAAGCGTATGTTCAAAA
 TACAATTGACAATCAACAAAGACTATCAACTGCTAATGCAATGCTGATAGAACCTCGTCAATATC
 AAAATCGCAGAGATACCACTCTCCCGATGCAAATTGAAACCATTAGGTTGGCATCAAGTAGCTACT
 AATGACCATTATGGACATGCAGTCGACAAGGGGATTTAATTGCCTATGCTTAGCTGGAAATTCAA
 AGGTTGGGATGCTTCCGTGTCAAATCCTAAAATGTTGTACACAAACAGCTCATTCCAACCAATCAA
 20 ATCAAAAAATCAATCGTGGACAAAATTATTGAAAGCTTAGTCGTAAGGGGGTTGACCAAAACAAA
 CGTGTCTGTTACCGTGTAACTCCATTGTAACGTAATGATACTGATTAGTTCCATTGCAATGCACCT
 AGAAGCTAAATCACAAGATGGCACATTAGAATTAAATGTTGCTATTCAAACACACAAGCATCATACA
 CTATGGATTATGCAACAGGAGAAATAACACTAAAT

25 **SEQ ID NO. 86**

MKRLHKLFIITVIATLGMLGVMTFGLPTQPQNVTPIVHADVNSSVDTSQEFQNNLKNAIGNLPFQYVNG
 IYEINNNQTNLNADVNVKAYVQNTIDNQQLSTANAMLDRTIRQYQNRDRTLDPANWKPLGWHQVAT
 NDHYGHAVDKGHLIAYALAGNFKGWDASVSNPQNVVTQTAHSNQSNQKINRGQNYYESLVRKAVDQNK
 30 RVRVTPLYRNDLVPFAMHLEAKSQDGTLEFNVAIPNTQASYTMDYATGEITLN

GBS 67

The following offers examples of preferred GBS 67 fragments. Nucleotide and amino acid sequence of GBS 67 sequences from serotype V isolated strain 2603 are set forth below as SEQ ID NOS: 87 and 88.

35 **SEQ ID NO: 87**

ATGAGAAAATACCAAAAATTTCTAAAATATTGACGTTAACGTCTTTTGTTGTCGCAAATACCGCT
 TAAATACCAATGTTAGGGAAAGTACCGTACCGGAAAATGGTCTAAAGGAAAGTTAGTTGTTAAAA
 AGACAGATGACCAGAACACACTTCAAAAGCTACCTTGTTAAAAACTACTGCTCATCCAGAA
 AGTAAAATAGAAAAAGTAACGTGAGCTAACAGGTGAAGCTACTTTGATAATCTCATACCTGGAGA
 40 TTAATACCTTATCAGAAGAACAGCGCCCGAAGGTTATAAAAAGACTAACAGACTTGGCAAGTTAAGG
 TTGAGAGTAATGGAAAAACTACGATACAAAATAGTGGTATAAAAATTCCACAATTGGACAAAATCAG
 GAAAGAAGTACGATACAGTATCCCCCAGGAATTATGAAGAGTACAAAGGAATCTTATAAAACTTGA
 GCACTGTTAAAGGTTCAAGTCCAAATGGAAAGTCAGAGGCAAAAGCAGTTAACCCATATTCAAGTGAAG
 GTGAGCATATAAGAGAAATTCCAGAGGGAACATTATCTAACACGTATTTCAGAAGTAGGTGATTTAGCT
 45 CATAATAAAATATAAAATTGAGTTAACGTGTCAGTGGAAAAACCATAGTAAAACCAAGTGGACAAACAAA
 GCCGTTAGATGTTGCTTCGTACTCGATAATTCTAACCTAACGATGGCCAAATTTCAA
 GGCATAATAAGCCAAGAAAGCTGCCAGTGGACCGCAGTAAAGATAATTAGGAGCAAAC

AGTGATAATAGGGTTGCATTAGTTACCTATGGTCAGATATTTGATGGTAGGAGTGTAGATGTCGT
 AAAAGGATTAAAGAAGATGATAAATATTATGGCCTCAAACATAAGTCACAATTAGACAGAGAATT
 ATAGTCATAAACAAATAACAAATAATGCTGAAGAGATTATAAAAAGGATTCCGACAGAACAGCTCTAAA
 GCTAAGTGGGATCTACTACCAATGGATTAACCTCAGAGCAACAAAGGAGTACTATCTTAGTAAAGT
 5 AGGAGAAACATTACTATGAAAGCCTCATGGAGGCAGATGATATTGAGTCAGTAAATCGAAATA
 GTCAAAAAATTATTGTCATGTAACGTGTTCTACGAGATCATATGCTTAAATAATTAA
 CTGGGTGCATCATATGAAAGCCAATTGAACAAATGAAAAAAATGGATATCTAAATAAAAGTAATT
 TCTACTTACTGATAAGCCGAGGATATAAAAGGAAATGGGAGAGTTACTTTGTTCCCTAGATA
 10 GTTATCAAACACAGATAATCTGGAAACTTACAAAACCTTACATTATTTAGATTAAATCTTAAATTAC
 CCTAAAGGTACAATTATCGAAATGGACCAGTGAAGAACATGGAACACCAACCAAACCTTATATAAA
 TAGTTAAAACAGAAAAATTATGACATTAAATTGGTATCGATATATCTGGTTTAGACAAGTT
 ATAATGAGGAGTATAAGAAAATCAAGATGGTACTTTCAAAAATTGAAAGAGGAAGCTTTAAACCT
 15 TCAGATGGAGAAATCACAGAACTAATGAGGTGTTCTTCCAAACCTGAGTACTACACCCCTATCGT
 AACCTCAGCCGATACATCTAACAAATGAAATTCTAAATTCAAGAACAAATTGAAACGATTAA
 CAAAAGAAAACACTCAATTGTAATGAACTATCGAAGATCCTATGGGTGATAAAATCAATTACAGCTT
 GGTAAATGGACAAACATTACAGCCAAGTGATTATACTTACAGGAAATGATGAAAGTGTAAATGAAGGA
 TGGTATTGCAACTGGTGGGCTAATAATGATGGTGAATACTAAGGGGTTAAATTAGAATACATCG
 GAAATAAAACTCTATGTTAGAGGTTGAATTAGGAGAAGGTCAAAAGTAACACTCACATATGATGTG
 20 AAAACTAGATGACAGTTATAAGTAACAAATTCTATGACACTAATGGTAGAACAAACATTGAATCCTAA
 GTCAGAGGATCCTAACACTAGAGATTTCATCCCTAAATTCTGATGTGAGAGAATATCCTA
 CAATAACGATTAACGAGAAGAAGTTAGGTGAAATTGAATTATAAAAGTGTAAAGATAATAAT
 AAGTTGCTCTCAAAGGAGCTACGTTGAACCTCAAGAATTAAATGAAAGATTATAAAACTTTATTAC
 25 AATAAAAAATAATAATTCAAAGTAGTAGTGACGGGAGAAACGGCAAATTCTACAAAGATTGAAAG
 ATGGCAAATATCAGTTAATAGAACAGTTGCGCCGGAGGATTATCAAAAAATTACTAATAAAACCAATT
 TTAACTTTGAAGTGGTAAAGGATCGATAAAAATATAATAGCTGTTAATAACAGATTCTGAATA
 TCATGAGGAAGGTGACAAGCATTAAATTACCAACACGCATTCCACCAAAGGAATTATTCTATGA
 CAGGTGGGAAAGGAATTCTATCTTCAATTAGGTGGAGCTATGATGCTATTGCAGGTGGAATT
 30 TATATTGGAAAAGGTATAAGAAATCTAGTGATATGTCCATCAAAAGAT
35 SEQ ID NO: 88
 MRKYQKFSKILTLSFLCSQIPLNTNVILGESTVPENGAKGKLVVKKTDDQNPLSKATFVLKTTAHE
 SKIEKVTAELTGEATFDNLIPGDYTLSEETAPEGYKKTNTQWQVKVESNGKTTIQNSGDKNSTIGQNZ
 EELDKQYPPTGIYEDTKESYKLEHVKGSPNGKSEAKVNPSSEGEHIREIPEGTLSKRISEVGDLA
 HNKYKIELTVSGKTIKVPDFQKPLDVVFVLDNSNSMNNDGPNFQRHNAKAKAAEALGTAVKDILGAN
 SDNRVALVTYGSDFDGRSVDVVKGFKEDDKYYGLQTKFTIQTENYHKQLTNNAEEIIKRIPTEAPK
 40 AKWGSTTNGLTPEQQKEYYLSKVGGETFTMKAFMEADDILSQVRNSQKIIIVHVTDGVPTRSYAINNFK
 LGASYESQFEQMKKNGYLNKSNFLTDKPEDIKGNGESYFLFPLDSYQTQIISGNLQKLHYLDLNLY
 PKGTIYRNGPVKEHGTPTKLYINSLQKNYDIFNFGIDISGFRQVYNEEYKKNQDGTFQKLKEEAFKL
 SDGEITELMRSFSSKPEYYTPIVTSADTSNNEILSKIQQQFETILTKEINSIVNGTIEDPMGDKINLQL
 GNGQTLQPSDYLQGDGSVMKDGIATGGPNNDGGILKGVKLEYIGNKLYVRGLNLGEQKVTLTYDV
 KLDSDFISNKFYDTNGRTTLNPKSEDPTLDFPIP KIRDVREYPTITIKNEKKLGEIEFIKVDKDNN
 KLLLKGATFELQEFNEDYKLYPIKNNNSKVVTGENGKISYKDLKDGYQQLIEAVSPEDYQKITNKPI
 LTFEVVKGSIKNIIAVNKQI~~SEYHEEGDKHLITNTHIPPKGI~~IPMTGGKILSFILIGGAMMSIAGGI
 45

GBS 67 contains a C-terminus transmembrane region which is indicated by the underlined region closest to the C-terminus of SEQ ID NO: 88 above. In one embodiment, one or more amino acids from the transmembrane region is removed and or the amino acid is truncated before the transmembrane region. An example of such a GBS 67 fragment is set forth below as SEQ ID NO: 89.

SEQ ID NO: 89

5 MRKYQKFSKILTSLFCLSQIPLNTNVLGESTVPENGAKGKLVVKKTDDQNKPLSKATFVLKTTAHPE
 SKIEKVTAELTGEATFDNLIPGDYTLSEETAPEGYKKTNTQWQVKVESNGKTTIQNSGDKNSTIGQNZ
 EELDKQYPPPTGIYEDTKESYKLEHVKGSPNGKSEAKAVNPYSSEGEHIREIPEGTLSKRISEVGDLA
 HNKYKIELTSGKTIVKPVDKQKPLDVFVLDSNSNSMNNNDGPNFQRHNAKAKAAEALGTAVKDILGAN
 SDNRVALVTYGSDIFDGRSDVVKGFKEDDKYYGLQTKFTIQTENYSHKQLTNNAEEIIKRIPTEAPK
 AKWGSTTNGLTPEQQKEYYLSKVGGETFTMKAFCMEADDILSQVNRNSQKIIIVHVTDGVPTRSYAINNFK
 10 LGASYESQFEQMCKNGYLNKSNFLTDKPEDIKGNGESYFLFPLDSYQTQIISGNLQKLHYLDLNLNY
 PKGTIYRNGPVKEHGTPTKLYINSLKQKNYDIFNFGIDISGFRQVYNEEYKKNQDGTQKLKEEAFKL
 SDGEITELMRSFSSKPEYYTPIVTSADTSNNEILSKIQQQFETILTKEINSIVNGTIEDPMGDKINLQL
 GNGQTLQPSDYTLQGNDGSVMKDGIAATGGPNNDGGILKGVKLEYIGNKLYVRGLNLGEQKVTLYDV
 KLDASFISNKFYDTNGRTTLNPKSEDPTLRFPIPKIRDVREYPTITIKNEKKLGEIEFIKVDKDNN
 KLLKGATFELQEFNEDYKLYLPIKNNNSKVVTGENGKISYKDLKGKYQLIEAVSPEDYQKITNKPI
 15 LTFEVVKGSIKNIIAVNKQISEYHEEGDKHLITNTHIPPKGIIPMTGGKGILS

GBS 67 contains an amino acid motif indicative of a cell wall anchor (an LPXTG motif):

20 **SEQ ID NO: 90** *I* PMTG. (shown in italics in SEQ ID NO: 88 above). In some recombinant host cell systems, it may be preferable to remove this motif to facilitate secretion of a recombinant GBS 67 protein from the host cell. Accordingly, in one preferred fragment of GBS 67 for use in the invention, the transmembrane and the cell wall anchor motif are removed from GBS 67. An example of such a GBS 67 fragment is set forth below as SEQ ID NO: 91.

SEQ ID NO: 91

25 MRKYQKFSKILTSLFCLSQIPLNTNVLGESTVPENGAKGKLVVKKTDDQNKPLSKATFVLKTTAHPE
 SKIEKVTAELTGEATFDNLIPGDYTLSEETAPEGYKKTNTQWQVKVESNGKTTIQNSGDKNSTIGQNZ
 EELDKQYPPPTGIYEDTKESYKLEHVKGSPNGKSEAKAVNPYSSEGEHIREIPEGTLSKRISEVGDLA
 HNKYKIELTSGKTIVKPVDKQKPLDVFVLDSNSNSMNNNDGPNFQRHNAKAKAAEALGTAVKDILGAN
 SDNRVALVTYGSDIFDGRSDVVKGFKEDDKYYGLQTKFTIQTENYSHKQLTNNAEEIIKRIPTEAPK
 AKWGSTTNGLTPEQQKEYYLSKVGGETFTMKAFCMEADDILSQVNRNSQKIIIVHVTDGVPTRSYAINNFK
 30 LGASYESQFEQMCKNGYLNKSNFLTDKPEDIKGNGESYFLFPLDSYQTQIISGNLQKLHYLDLNLNY
 PKGTIYRNGPVKEHGTPTKLYINSLKQKNYDIFNFGIDISGFRQVYNEEYKKNQDGTQKLKEEAFKL
 SDGEITELMRSFSSKPEYYTPIVTSADTSNNEILSKIQQQFETILTKEINSIVNGTIEDPMGDKINLQL
 GNGQTLQPSDYTLQGNDGSVMKDGIAATGGPNNDGGILKGVKLEYIGNKLYVRGLNLGEQKVTLYDV
 KLDASFISNKFYDTNGRTTLNPKSEDPTLRFPIPKIRDVREYPTITIKNEKKLGEIEFIKVDKDNN
 KLLKGATFELQEFNEDYKLYLPIKNNNSKVVTGENGKISYKDLKGKYQLIEAVSPEDYQKITNKPI
 35 LTFEVVKGSIKNIIAVNKQISEYHEEGDKHLITNTHIPPKGI

The compositions of the invention may also include combinations including one or more known GBS antigens in combination with GBS 80.

40 There is an upper limit to the number of GBS antigens which will be in the compositions of the invention. Preferably, the number of GBS antigens in a composition of the invention is less than 20, less than 19, less than 18, less than 17, less than 16, less than 15, less than 14, less than 13, less than 12, less than 11, less than 10, less than 9, less than 8, less than 7, less than 6, less than 5, less than 4, or less than 3. Still more preferably, the number of GBS antigens in a composition of the invention

is less than 6, less than 5, or less than 4. Still more preferably, the number of GBS antigens in a composition of the invention is 3.

The GBS antigens used in the invention are preferably isolated, i.e., separate and discrete, from the whole organism with which the molecule is found in nature or, when the polynucleotide or polypeptide is not found in nature, is sufficiently free of other biological macromolecules so that the polynucleotide or polypeptide can be used for its intended purpose.

Fusion Proteins

The GBS antigens used in the invention may be present in the composition as individual separate polypeptides, but it is preferred that at least two (i.e. 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, or 18) of the antigens are expressed as a single polypeptide chain (a "hybrid" or "fusion" polypeptide). Such fusion polypeptides offer two principal advantages: first, a polypeptide that may be unstable or poorly expressed on its own can be assisted by adding a suitable fusion partner that overcomes the problem; second, commercial manufacture is simplified as only one expression and purification need be employed in order to produce two polypeptides which are both antigenically useful.

The fusion polypeptide may comprise two or more polypeptide sequences from the group consisting of GBS 80, GBS 91, GBS 104, GBS 184, GBS 276, GBS 305, GBS 322, GBS 330, GBS 338, GBS 361, GBS 404, GBS 690 and GBS 691. Preferably, the polypeptide sequences are selected from the group consisting of GBS 80, GBS 104 and GBS 322. Most preferably, the fusion peptide includes a polypeptide sequence from GBS 80. Accordingly, the invention includes a fusion peptide comprising a first amino acid sequence and a second amino acid sequence, wherein said first and second amino acid sequences are selected from a GBS antigen or a fragment thereof of the above antigen group. Preferably, the first and second amino acid sequences in the fusion polypeptide comprise different epitopes.

EXAMPLE 7: Examples of fragments for fusion proteins from GBS 80 with GBS 104, and GBS 322

Examples of GBS fragments for fusion proteins are provided from GBS 322, GBS 104, and GBS 80. One example of a fragment of GBS 322 in a fusion protein is a 407 amino acid fragment with the signal peptide removed. Fragments of GBS 104 may also be incorporated in fusion proteins. An example of GBS 104 fragments includes an 830 amino acid fragment, a 359 amino acid fragment from near the N-terminus, a 581 amino acid fragment from near the N-terminus, and a 740 amino acid fragment from near the N-terminus. Examples of GBS 80 fragments include a 446 amino acid fragment and a 235 amino acid fragment. Table 13 below summarizes the examples of fragments for fusion proteins and their locations within the corresponding full length GBS protein.

Table 13: Active Maternal Immunization Assay using combinations of GBS 80 with GBS 104 and GBS 322

GBS	Size (AA)	SEQ ID NO	From ... to
322	407	92	25-432
104	830	96	28-858
104 N1	359	97	28-387
104 N2	581	98	28-609
104 N3	740	99	28-768
80	446	100	37-483
80N	235	101	37-272

5

Hybrids (or fusions) consisting of amino acid sequences from two, three, four, five, six, seven, eight, nine, or ten GBS antigens are preferred. In particular, hybrids consisting of amino acid sequences from two, three, four, or five GBS antigens are preferred.

Different hybrid polypeptides may be mixed together in a single formulation. Within such combinations, a GBS antigen may be present in more than one hybrid polypeptide and/or as a non-hybrid polypeptide. It is preferred, however, that an antigen is present either as a hybrid or as a non-hybrid, but not as both.

Hybrid polypeptides can be represented by the formula $\text{NH}_2\text{-A}\text{-}\{-\text{X-L-}\}_n\text{-B-COOH}$, wherein: X is an amino acid sequence of a GBS antigen or a fragment thereof from the antigen group set forth above; L is an optional linker amino acid sequence; A is an optional N-terminal amino acid sequence; B is an optional C-terminal amino acid sequence; and n is 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14 or 15.

If a -X- moiety has a leader peptide sequence in its wild-type form, this may be included or omitted in the hybrid protein. In some embodiments, the leader peptides will be deleted except for that of the -X- moiety located at the N-terminus of the hybrid protein *i.e.* the leader peptide of X₁ will be retained, but the leader peptides of X₂ ... X_n will be omitted. This is equivalent to deleting all leader peptides and using the leader peptide of X₁ as moiety -A-.

For each n instances of {-X-L-}, linker amino acid sequence -L- may be present or absent. For instance, when n=2 the hybrid may be $\text{NH}_2\text{-X}_1\text{-L}_1\text{-X}_2\text{-L}_2\text{-COOH}$, $\text{NH}_2\text{-X}_1\text{-X}_2\text{-COOH}$, $\text{NH}_2\text{-X}_1\text{-L}_1\text{-X}_2\text{-COOH}$, $\text{NH}_2\text{-X}_1\text{-X}_2\text{-L}_2\text{-COOH}$, *etc.* Linker amino acid sequence(s) -L- will typically be short (*e.g.* 20 or fewer amino acids *i.e.* 19, 18, 17, 16, 15, 14, 13, 12, 11, 10, 9, 8, 7, 6, 5, 4, 3, 2, 1). Examples comprise short peptide sequences which facilitate cloning, poly-glycine linkers (*i.e.* comprising Gly_n where n = 2, 3, 4, 5, 6, 7, 8, 9, 10 or more), and histidine tags (*i.e.* His_n where n = 3, 4, 5, 6, 7, 8, 9, 10 or more). Other suitable linker amino acid sequences will be apparent to those skilled in the art. A useful linker is GSGGGG, with the Gly-Ser dipeptide being formed from a

*Bam*HI restriction site, thus aiding cloning and manipulation, and the (Gly)₄ tetrapeptide being a typical poly-glycine linker.

-A- is an optional N-terminal amino acid sequence. This will typically be short (e.g. 40 or fewer amino acids *i.e.* 39, 38, 37, 36, 35, 34, 33, 32, 31, 30, 29, 28, 27, 26, 25, 24, 23, 22, 21, 20, 19, 5 18, 17, 16, 15, 14, 13, 12, 11, 10, 9, 8, 7, 6, 5, 4, 3, 2, 1). Examples include leader sequences to direct protein trafficking, or short peptide sequences which facilitate cloning or purification (e.g. histidine tags *i.e.* His_n where $n = 3, 4, 5, 6, 7, 8, 9, 10$ or more). Other suitable N-terminal amino acid sequences will be apparent to those skilled in the art. If X₁ lacks its own N-terminus methionine, -A- is preferably an oligopeptide (e.g. with 1, 2, 3, 4, 5, 6, 7 or 8 amino acids) which provides a 10 N-terminus methionine.

-B- is an optional C-terminal amino acid sequence. This will typically be short (e.g. 40 or fewer amino acids *i.e.* 39, 38, 37, 36, 35, 34, 33, 32, 31, 30, 29, 28, 27, 26, 25, 24, 23, 22, 21, 20, 19, 18, 17, 16, 15, 14, 13, 12, 11, 10, 9, 8, 7, 6, 5, 4, 3, 2, 1). Examples include sequences to direct protein trafficking, short peptide sequences which facilitate cloning or purification (e.g. comprising 15 histidine tags *i.e.* His_n where $n = 3, 4, 5, 6, 7, 8, 9, 10$ or more), or sequences which enhance protein stability. Other suitable C-terminal amino acid sequences will be apparent to those skilled in the art. Most preferably, n is 2 or 3.

20 **EXAMPLE 8: Active Maternal Immunization Assay using fusion proteins of**
Fragments of GBS 80, GBS 67, and GBS 322

In this example, fusion proteins of GBS antigens was used in the Active Maternal Immunization Assay with an isolate challenge of different GBS strains. In these experiments, the challenge dose for the different GBS strains was sufficient to kill approximately 70 – 90% of unimmunized pups and is equal to 10 x LD 50% (where LD 50% is the statistically derived Median 25 Lethal Dose). The maternal mice were immunized according to the Active Maternal Immunization Assay schedule described above with the fusion proteins of a GBS 80 antigen with GBS 322 antigen in the GBS strains set forth in Table 14 below. Survival % was observed with the GBS fusion proteins. As shown in Table 14, in this particular challenge study, the survival rates for the fusion proteins in all of the GBS strains achieved up to 79%.

Table 14: Active Maternal Immunization Assay using fusion proteins of GBS 80 with GBS 322

	COH1 (III)		CJB111 (V)		515 (Ia)		DK21 (II)		2603 (V)	
GBS	Dead/treated	Survival %								
80N-322	16/40	60	8/39	79	12/28	57	7/19	63	8/37	78
80	4/24	83								
PBS	35/40	12	27/35	23	32/39	18	31/40	22	33/40	17
80-322	12/27	55							12/38	68
80	0/33	100	28/40	30						
322									1/16	94
PBS	19/20	5	38/39	2	25/29	14			19/26	27

5

Nucleic Acids

The invention also provides nucleic acid encoding the GBS antigens and/or the hybrid fusion polypeptides of the invention. Furthermore, the invention provides nucleic acid which can hybridise to these nucleic acids, preferably under "high stringency" conditions (e.g. 65°C in a 0.1xSSC, 0.5% SDS solution).

Polypeptides of the invention can be prepared by various means (e.g. recombinant expression, purification from cell culture, chemical synthesis, etc.) and in various forms (e.g. native, fusions, non-glycosylated, lipidated, etc.). They are preferably prepared in substantially pure form (i.e. substantially free from other GAS or host cell proteins).

Nucleic acid according to the invention can be prepared in many ways (e.g. by chemical synthesis, from genomic or cDNA libraries, from the organism itself, etc.) and can take various forms (e.g. single stranded, double stranded, vectors, probes, etc.). They are preferably prepared in substantially pure form (i.e. substantially free from other GBS or host cell nucleic acids).

The term "nucleic acid" includes DNA and RNA, and also their analogues, such as those containing modified backbones (e.g. phosphorothioates, etc.), and also peptide nucleic acids (PNA), etc. The invention includes nucleic acid comprising sequences complementary to those described above (e.g. for antisense or probing purposes).

The invention also provides a process for producing a polypeptide of the invention, comprising the step of culturing a host cell transformed with nucleic acid of the invention under conditions which induce polypeptide expression.

The invention provides a process for producing a polypeptide of the invention, comprising the step of synthesising at least part of the polypeptide by chemical means.

The invention provides a process for producing nucleic acid of the invention, comprising the step of amplifying nucleic acid using a primer-based amplification method (e.g. PCR).

5 The invention provides a process for producing nucleic acid of the invention, comprising the step of synthesising at least part of the nucleic acid by chemical means.

Purification and Recombinant Expression

10 The GBS antigens of the invention may be isolated from *Streptococcus agalactiae*, or they may be recombinantly produced, for instance, in a heterologous host. Preferably, the GBS antigens are prepared using a heterologous host. The heterologous host may be prokaryotic (e.g. a bacterium) or eukaryotic. It is preferably *E. coli*, but other suitable hosts include *Bacillus subtilis*, *Vibrio cholerae*, *Salmonella typhi*, *Salmonella typhimurium*, *Neisseria lactamica*, *Neisseria cinerea*, *Mycobacteria* (e.g. *M. tuberculosis*), yeasts, etc.

15 Recombinant production of polypeptides is facilitated by adding a tag protein to the GBS antigen to be expressed as a fusion protein comprising the tag protein and the GBS antigen. Such tag proteins can facilitate purification, detection and stability of the expressed protein. Tag proteins suitable for use in the invention include a polyarginine tag (Arg-tag), polyhistidine tag (His-tag), FLAG-tag, Strep-tag, c-myc-tag, S-tag, calmodulin-binding peptide, cellulose-binding domain, SBP-
20 tag,, chitin-binding domain, glutathione S-transferase-tag (GST), maltose-binding protein, transcription termination anti-terminant factor (NusA), *E. coli* thioredoxin (TrxA) and protein disulfide isomerase I (DsbA). Preferred tag proteins include His-tag and GST. A full discussion on the use of tag proteins can be found at Ref. 3.

25 After purification, the tag proteins may optionally be removed from the expressed fusion protein, i.e., by specifically tailored enzymatic treatments known in the art. Commonly used proteases include enterokinase, tobacco etch virus (TEV), thrombin, and factor X_a.

GBS polysaccharides

30 The compositions of the invention may be further improved by including GBS polysaccharides. Preferably, the GBS antigen and the saccharide each contribute to the immunological response in a recipient. The combination is particularly advantageous where the saccharide and polypeptide provide protection from different GBS serotypes.

35 The combined antigens may be present as a simple combination where separate saccharide and polypeptide antigens are administered together, or they may be present as a conjugated combination, where the saccharide and polypeptide antigens are covalently linked to each other.

Thus the invention provides an immunogenic composition comprising (i) one or more GBS polypeptide antigens and (ii) one or more GBS saccharide antigens. The polypeptide and the polysaccharide may advantageously be covalently linked to each other to form a conjugate.

Between them, the combined polypeptide and saccharide antigens preferably cover (or provide protection from) two or more GBS serotypes (e.g. 2, 3, 4, 5, 6, 7, 8 or more serotypes). The serotypes of the polypeptide and saccharide antigens may or may not overlap. For example, the polypeptide might protect against serogroup II or V, while the saccharide protects against either serogroups Ia, Ib, or III. Preferred combinations protect against the following groups of serotypes: (1) serotypes Ia and Ib, (2) serotypes Ia and II, (3) serotypes Ia and III, (4) serotypes Ia and IV, (5) serotypes Ia and V, (6) serotypes Ia and VI, (7) serotypes Ia and VII, (8) serotypes Ia and VIII, (9) serotypes Ib and II, (10) serotypes Ib and III, (11) serotypes Ib and IV, (12) serotypes Ib and V, (13) serotypes Ib and VI, (14) serotypes Ib and VII, (15) serotypes Ib and VIII, (16) serotypes II and III, (17) serotypes II and IV, (18) serotypes II and V, (19) serotypes II and VI, (20) serotypes II and VII, (21) serotypes II and VIII, (22) serotypes III and IV, (23) serotypes III and V, (24) serotypes III and VI, (25) serotypes III and VII, (26) serotypes III and VIII, (27) serotypes IV and V, (28) serotypes IV and VI, (29) serotypes IV and VII, (30) serotypes IV and VIII, (31) serotypes V and VI, (32) serotypes V and VII, (33) serotypes V and VIII, (34) serotypes VI and VII, (35) serotypes VI and VIII, and (36) serotypes VII and VIII.

Still more preferably, the combinations protect against the following groups of serotypes: (1) serotypes Ia and II, (2) serotypes Ia and V, (3) serotypes Ib and II, (4) serotypes Ib and V, (5) serotypes III and II, and (6) serotypes III and V. Most preferably, the combinations protect against serotypes III and V.

Protection against serotypes II and V is preferably provided by polypeptide antigens.

Protection against serotypes Ia, Ib and/or III may be polypeptide or saccharide antigens.

In one embodiment, the immunogenic composition comprises a GBS saccharide antigen and at least two GBS polypeptide antigens or fragments thereof, wherein said GBS saccharide antigen comprises a saccharide selected from GBS serotype Ia, Ib, and III, and wherein said GBS polypeptide antigens comprise a combination of at least two polypeptide or a fragment thereof selected from the antigen group consisting of GBS 80, GBS 91, GBS 104, GBS 184, GBS 276, GBS 305, GBS 322, GBS 330, GBS 338, GBS 361, GBS 404, GBS 690, and GBS 691. Preferably, the combination includes one or more of GBS 80, GBS 104 and GBS 322. Still more preferably, the combination includes GBS 80 or a fragment thereof.

In certain embodiments, the compositions of the invention do not include a GBS polysaccharide. In certain embodiments, the combination does not include one or more of the GBS antigens selected from the group consisting of GBS 4, GBS 22, GBS 85, GBS 338 and GBS 361.

Immunogenic compositions and medicaments

Compositions of the invention are preferably immunogenic compositions, and are more preferably vaccine compositions. The pH of the composition is preferably between 6 and 8, preferably about 7. The pH may be maintained by the use of a buffer. The composition may be 5 sterile and/or pyrogen-free. The composition may be isotonic with respect to humans.

Vaccines according to the invention may either be prophylactic (*i.e.* to prevent infection) or therapeutic (*i.e.* to treat infection), but will typically be prophylactic. Accordingly, the invention includes a method for the therapeutic or prophylactic treatment of a *Streptococcus agalactiae* 10 infection in an animal susceptible to streptococcal infection comprising administering to said animal a therapeutic or prophylactic amount of the immunogenic compositions of the invention.

The invention also provides a composition of the invention for use as a medicament. The medicament is preferably able to raise an immune response in a mammal (*i.e.* it is an immunogenic composition) and is more preferably a vaccine.

The invention also provides the use of the compositions of the invention in the manufacture of 15 a medicament for raising an immune response in a mammal. The medicament is preferably a vaccine.

The invention also provides for a kit comprising a first component comprising a combination of GBS antigens.

The invention also provides a delivery device pre-filled with the immunogenic compositions of the invention.

20 The invention also provides a method for raising an immune response in a mammal comprising the step of administering an effective amount of a composition of the invention. The immune response is preferably protective and preferably involves antibodies and/or cell-mediated immunity. The method may raise a booster response.

The mammal is preferably a human. Where the vaccine is for prophylactic use, the human is 25 preferably a female (either of child bearing age or a teenager). Alternatively, the human may be elderly (e.g., over the age of 50, 55, 60, 65, 70 or 75) and may have an underlying disease such as diabetes or cancer. Where the vaccine is for therapeutic use, the human is preferably a pregnant female or an elderly adult.

These uses and methods are preferably for the prevention and/or treatment of a disease caused 30 by *Streptococcus agalactiae*. The compositions may also be effective against other streptococcal bacteria.

One way of checking efficacy of therapeutic treatment involves monitoring GBS infection after administration of the composition of the invention. One way of checking efficacy of prophylactic treatment involves monitoring immune responses against the GBS antigens in the 35 compositions of the invention after administration of the composition.

Compositions of the invention will generally be administered directly to a patient. Direct delivery may be accomplished by parenteral injection (e.g. subcutaneously, intraperitoneally, intradermally, intravenously, intramuscularly, or to the interstitial space of a tissue), or by rectal, oral (e.g. tablet, spray), vaginal, topical, transdermal {e.g. see ref. 4} or transcutaneous {e.g. see refs. 5 & 6}, intranasal {e.g. see ref. 7}, ocular, aural, pulmonary or other mucosal administration.

5 The invention may be used to elicit systemic and/or mucosal immunity.

Dosage treatment can be a single dose schedule or a multiple dose schedule. Multiple doses may be used in a primary immunisation schedule and/or in a booster immunisation schedule. In a 10 multiple dose schedule the various doses may be given by the same or different routes e.g. a parenteral prime and mucosal boost, a mucosal prime and parenteral boost, etc.

The compositions of the invention may be prepared in various forms. For example, the 15 compositions may be prepared as injectables, either as liquid solutions or suspensions. Solid forms suitable for solution in, or suspension in, liquid vehicles prior to injection can also be prepared (e.g. a lyophilised composition). The composition may be prepared for topical administration e.g. as an ointment, cream or powder. The composition may be prepared for oral administration e.g. as a tablet or capsule, as a spray, or as a syrup (optionally flavoured). The composition may be prepared for pulmonary administration e.g. as an inhaler, using a fine powder or a spray. The composition may be prepared as a suppository or pessary. The composition may be prepared for nasal, aural or ocular 20 administration e.g. as drops. The composition may be in kit form, designed such that a combined composition is reconstituted just prior to administration to a patient. Such kits may comprise one or more antigens in liquid form and one or more lyophilised antigens.

25 Immunogenic compositions used as vaccines comprise an immunologically effective amount of antigen(s), as well as any other components, as needed. By 'immunologically effective amount', it is meant that the administration of that amount to an individual, either in a single dose or as part of a series, is effective for treatment or prevention. This amount varies depending upon the health and physical condition of the individual to be treated, age, the taxonomic group of individual to be treated (e.g. non-human primate, primate, etc.), the capacity of the individual's immune system to synthesise 30 antibodies, the degree of protection desired, the formulation of the vaccine, the treating doctor's assessment of the medical situation, and other relevant factors. It is expected that the amount will fall in a relatively broad range that can be determined through routine trials.

Further Components of the Composition

The composition of the invention will typically, in addition to the components mentioned above, comprise one or more 'pharmaceutically acceptable carriers', which include any carrier that 35 does not itself induce the production of antibodies harmful to the individual receiving the

composition. Suitable carriers are typically large, slowly metabolised macromolecules such as proteins, polysaccharides, polylactic acids, polyglycolic acids, polymeric amino acids, amino acid copolymers, and lipid aggregates (such as oil droplets or liposomes). Such carriers are well known to those of ordinary skill in the art. The vaccines may also contain diluents, such as water, saline, 5 glycerol, *etc.* Additionally, auxiliary substances, such as wetting or emulsifying agents, pH buffering substances, and the like, may be present. A thorough discussion of pharmaceutically acceptable excipients is available in reference 8.

Vaccines of the invention may be administered in conjunction with other immunoregulatory agents. In particular, compositions will usually include an adjuvant.

10 Preferred further adjuvants include, but are not limited to, one or more of the following set forth below:

A. Mineral Containing Compositions

Mineral containing compositions suitable for use as adjuvants in the invention include mineral salts, such as aluminium salts and calcium salts. The invention includes mineral salts such as 15 hydroxides (*e.g.* oxyhydroxides), phosphates (*e.g.* hydroxyphosphates, orthophosphates), sulphates, *etc.* {*e.g.* see chapters 8 & 9 of ref. 9}), or mixtures of different mineral compounds, with the compounds taking any suitable form (*e.g.* gel, crystalline, amorphous, *etc.*), and with adsorption being preferred. The mineral containing compositions may also be formulated as a particle of metal salt. See ref. 10.

20 B. Oil-Emulsions

Oil-emulsion compositions suitable for use as adjuvants in the invention include squalene-water emulsions, such as MF59 (5% Squalene, 0.5% Tween 80, and 0.5% Span 85, formulated into submicron particles using a microfluidizer). See WO90/14837. See also, Frey et al., "Comparison of the safety, tolerability, and immunogenicity of a MF59-adjuvanted influenza 25 vaccine and a non-adjuvanted influenza vaccine in non-elderly adults", Vaccine (2003) 21:4234–4237.

Particularly preferred adjuvants for use in the compositions are submicron oil-inwater emulsions. Preferred submicron oil-in-water emulsions for use herein are squalene/water emulsions optionally containing varying amounts of MTP-PE, such as a submicron oil-in-water 30 emulsion containing 4-5% w/v squalene, 0.25-1.0% w/v Tween 80™ (polyoxyethylenglycosorbitan monooleate), and/or 0.25-1.0% Span 85™ (sorbitan trioleate), and, optionally, N-acetylmuramyl-L-alanyl-D-isoglutaminyl-L-alanine-2-(1'-2'-dipalmitoyl-*sn*-glycero-3-hydroxyphosphoryloxy)-ethylamine (MTP-PE), for example, the submicron oil-in-water emulsion known as "MF59" (International Publication No. WO 90/14837; U.S. Patent Nos. 35 6,299,884 and 6,451,325, incorporated herein by reference in their entireties; and Ott et al.,

"MF59 -- Design and Evaluation of a Safe and Potent Adjuvant for Human Vaccines" in *Vaccine Design: The Subunit and Adjuvant Approach* (Powell, M.F. and Newman, M.J. eds.) Plenum Press, New York, 1995, pp. 277-296). MF59 contains 4-5% w/v Squalene (e.g., 4.3%), 0.25-0.5% w/v Tween 80TM, and 0.5% w/v Span 85TM and optionally contains various amounts of

5 MTP-PE, formulated into submicron particles using a microfluidizer such as Model 110Y microfluidizer (Microfluidics, Newton, MA). For example, MTP-PE may be present in an amount of about 0-500 µg/dose, more preferably 0-250 µg/dose and most preferably, 0-100 µg/dose. As used herein, the term "MF59-0" refers to the above submicron oil-in-water emulsion lacking MTP-PE, while the term MF59-MTP denotes a formulation that contains MTP-PE. For

10 instance, "MF59-100" contains 100 µg MTP-PE per dose, and so on. MF69, another submicron oil-in-water emulsion for use herein, contains 4.3% w/v squalene, 0.25% w/v Tween 80TM, and 0.75% w/v Span 85TM and optionally MTP-PE. Yet another submicron oil-in-water emulsion is MF75, also known as SAF, containing 10% squalene, 0.4% Tween 80TM, 5% pluronic-blocked polymer L121, and thr-MDP, also microfluidized into a submicron emulsion. MF75-MTP

15 denotes an MF75 formulation that includes MTP, such as from 100-400 µg MTP-PE per dose.

Submicron oil-in-water emulsions, methods of making the same and immunostimulating agents, such as muramyl peptides, for use in the compositions, are described in detail in International Publication No. WO 90114837 and U.S. Patent Nos. 6,299,884 and 6,451,325, incorporated herein by reference in their entireties.

20 Complete Freund's adjuvant (CFA) and incomplete Freund's adjuvant (IFA) may also be used as adjuvants in the invention.

C. Saponin Formulations

25 Saponin formulations, may also be used as adjuvants in the invention. Saponins are a heterologous group of sterol glycosides and triterpenoid glycosides that are found in the bark, leaves, stems, roots and even flowers of a wide range of plant species. Saponin from the bark of the *Quillaia saponaria* Molina tree have been widely studied as adjuvants. Saponin can also be commercially obtained from *Smilax ornata* (sarsaparilla), *Gypsophilla paniculata* (brides veil), and *Saponaria officianalis* (soap root). Saponin adjuvant formulations include purified formulations, such as QS21, as well as lipid formulations, such as ISCOMs.

30 Saponin compositions have been purified using High Performance Thin Layer Chromatography (HP-LC) and Reversed Phase High Performance Liquid Chromatography (RP-HPLC). Specific purified fractions using these techniques have been identified, including QS7, QS17, QS18, QS21, QH-A, QH-B and QH-C. Preferably, the saponin is QS21. A method of production of

QS21 is disclosed in U.S. Patent No. 5,057,540. Saponin formulations may also comprise a sterol, such as cholesterol (see WO 96/33739).

Combinations of saponins and cholesterol can be used to form unique particles called Immunostimulating Complexes (ISCOMs). ISCOMs typically also include a phospholipid such as 5 phosphatidylethanolamine or phosphatidylcholine. Any known saponin can be used in ISCOMs. Preferably, the ISCOM includes one or more of Quil A, QHA and QHC. ISCOMs are further described in EP 0 109 942, WO 96/11711 and WO 96/33739. Optionally, the ISCOMS may be devoid of additional detergent. See ref. 11.

A review of the development of saponin based adjuvants can be found at ref. 12.

10 C. Virosomes and Virus Like Particles (VLPs)

Virosomes and Virus Like Particles (VLPs) can also be used as adjuvants in the invention. These structures generally contain one or more proteins from a virus optionally combined or 15 formulated with a phospholipid. They are generally non-pathogenic, non-replicating and generally do not contain any of the native viral genome. The viral proteins may be recombinantly produced or isolated from whole viruses. These viral proteins suitable for use in virosomes or VLPs include 20 proteins derived from influenza virus (such as HA or NA), Hepatitis B virus (such as core or capsid proteins), Hepatitis E virus, measles virus, Sindbis virus, Rotavirus, Foot-and-Mouth Disease virus, Retrovirus, Norwalk virus, human Papilloma virus, HIV, RNA-phages, Q β -phage (such as coat proteins), GA-phage, fr-phage, AP205 phage, and Ty (such as retrotransposon Ty protein p1). VLPs are discussed further in WO 03/024480, WO 03/024481, and Refs. 13, 14, 15 and 16. Virosomes are discussed further in, for example, Ref. 17

D. Bacterial or Microbial Derivatives

Adjuvants suitable for use in the invention include bacterial or microbial derivatives such as:

(1) *Non-toxic derivatives of enterobacterial lipopolysaccharide (LPS)*

25 Such derivatives include Monophosphoryl lipid A (MPL) and 3-O-deacylated MPL (3dMPL). 3dMPL is a mixture of 3 De-O-acylated monophosphoryl lipid A with 4, 5 or 6 acylated chains. A preferred "small particle" form of 3 De-O-acylated monophosphoryl lipid A is disclosed in EP 0 689 454. Such "small particles" of 3dMPL are small enough to be sterile filtered through a 0.22 micron 30 membrane (see EP 0 689 454). Other non-toxic LPS derivatives include monophosphoryl lipid A mimics, such as aminoalkyl glucosaminide phosphate derivatives e.g. RC-529. See Ref. 18.

(2) *Lipid A Derivatives*

Lipid A derivatives include derivatives of lipid A from *Escherichia coli* such as OM-174. OM-174 is described for example in Ref. 19 and 20.

5 (3) *Immunostimulatory oligonucleotides*

Immunostimulatory oligonucleotides suitable for use as adjuvants in the invention include nucleotide sequences containing a CpG motif (a sequence containing an unmethylated cytosine followed by guanosine and linked by a phosphate bond). Bacterial double stranded RNA or oligonucleotides containing palindromic or poly(dG) sequences have also been shown to be immunostimulatory.

10 The CpG's can include nucleotide modifications/analogs such as phosphorothioate modifications and can be double-stranded or single-stranded. Optionally, the guanosine may be replaced with an analog such as 2'-deoxy-7-deazaguanosine. See ref. 21, WO 02/26757 and WO 99/62923 for examples of possible analog substitutions. The adjuvant effect of CpG oligonucleotides is further discussed in Refs. 22, 23, WO 98/40100, U.S. Patent No. 6,207,646, U.S. Patent No. 6,239,116, and U.S. Patent No. 6,429,199.

15 The CpG sequence may be directed to TLR9, such as the motif GTCGTT or TTTCGTT. See ref. 24. The CpG sequence may be specific for inducing a Th1 immune response, such as a CpG-A ODN, or it may be more specific for inducing a B cell response, such a CpG-B ODN. CpG-A and CpG-B ODNs are discussed in refs. 25, 26 and WO 01/95935. Preferably, the CpG is a CpG-A ODN.

20 Preferably, the CpG oligonucleotide is constructed so that the 5' end is accessible for receptor recognition. Optionally, two CpG oligonucleotide sequences may be attached at their 3' ends to form "immunomers". See, for example, refs. 27, 28, 29 and WO 03/035836.

(4) *ADP-ribosylating toxins and detoxified derivatives thereof.*

25 Bacterial ADP-ribosylating toxins and detoxified derivatives thereof may be used as adjuvants in the invention. Preferably, the protein is derived from *E. coli* (i.e., *E. coli* heat labile enterotoxin ("LT"), cholera ("CT"), or pertussis ("PT"). The use of detoxified ADP-ribosylating toxins as mucosal adjuvants is described in WO 95/17211 and as parenteral adjuvants in WO 98/42375. Preferably, the adjuvant is a detoxified LT mutant such as LT-K63.

E. Human Immunomodulators

30 Human immunomodulators suitable for use as adjuvants in the invention include cytokines, such as interleukins (e.g. IL-1, IL-2, IL-4, IL-5, IL-6, IL-7, IL-12, etc.), interferons (e.g. interferon- γ), macrophage colony stimulating factor, and tumor necrosis factor.

F. Bioadhesives and Mucoadhesives

Bioadhesives and mucoadhesives may also be used as adjuvants in the invention. Suitable bioadhesives include esterified hyaluronic acid microspheres (Ref. 30) or mucoadhesives such as cross-linked derivatives of poly(acrylic acid), polyvinyl alcohol, polyvinyl pyrrolidone,

polysaccharides and carboxymethylcellulose. Chitosan and derivatives thereof may also be used as adjuvants in the invention. E.g., ref. 31.

G. Microparticles

5 Microparticles may also be used as adjuvants in the invention. Microparticles (*i.e.* a particle of ~100nm to ~150μm in diameter, more preferably ~200nm to ~30μm in diameter, and most preferably ~500nm to ~10μm in diameter) formed from materials that are biodegradable and non-toxic (*e.g.* a poly(α -hydroxy acid), a polyhydroxybutyric acid, a polyorthoester, a polyanhydride, a polycaprolactone, *etc.*), with poly(lactide-co-glycolide) are preferred, optionally treated to have a negatively-charged surface (*e.g.* with SDS) or a positively-charged surface (*e.g.* with a cationic 10 detergent, such as CTAB).

H. Liposomes

Examples of liposome formulations suitable for use as adjuvants are described in U.S. Patent No. 6,090,406, U.S. Patent No. 5,916,588, and EP 0 626 169.

I. Polyoxyethylene ether and Polyoxyethylene Ester Formulations

15 Adjuvants suitable for use in the invention include polyoxyethylene ethers and polyoxyethylene esters. Ref. 32. Such formulations further include polyoxyethylene sorbitan ester surfactants in combination with an octoxynol (Ref. 33) as well as polyoxyethylene alkyl ethers or ester surfactants in combination with at least one additional non-ionic surfactant such as an octoxynol (Ref. 34).

20 Preferred polyoxyethylene ethers are selected from the following group: polyoxyethylene-9-lauryl ether (laureth 9), polyoxyethylene-9-stearyl ether, polyoxyethylene-8-stearyl ether, polyoxyethylene-4-lauryl ether, polyoxyethylene-35-lauryl ether, and polyoxyethylene-23-lauryl ether.

J. Polyphosphazene (PCPP)

25 PCPP formulations are described, for example, in Ref. 35 and 36.

K. Muramyl peptides

Examples of muramyl peptides suitable for use as adjuvants in the invention include N-acetyl-muramyl-L-threonyl-D-isoglutamine (thr-MDP), N-acetyl-normuramyl-L-alanyl-D-isoglutamine (nor-MDP), and N-acetylmuramyl-L-alanyl-D-isoglutaminyl-L-alanine-2-(1'-2'-dipalmitoyl-*sn*-glycero-3'-30 hydroxyphosphoryloxy)-ethylamine MTP-PE).

L. Imidazoquinolone Compounds.

Examples of imidazoquinolone compounds suitable for use adjuvants in the invention include Imiquamod and its homologues, described further in Ref. 37 and 38.

The invention may also comprise combinations of aspects of one or more of the adjuvants identified above. For example, the following adjuvant compositions may be used in the invention:

- (1) a saponin and an oil-in-water emulsion (ref. 39);
- 5 (2) a saponin (e.g., QS21) + a non-toxic LPS derivative (e.g., 3dMPL) (see WO 94/00153);
- (3) a saponin (e.g., QS21) + a non-toxic LPS derivative (e.g., 3dMPL) + a cholesterol;
- (4) a saponin (e.g. QS21) + 3dMPL + IL-12 (optionally + a sterol) (Ref. 40);
- (5) combinations of 3dMPL with, for example, QS21 and/or oil-in-water emulsions (Ref. 10 41);
- (6) SAF, containing 10% Squalane, 0.4% Tween 80, 5% pluronic-block polymer L121, and thr-MDP, either microfluidized into a submicron emulsion or vortexed to generate a larger particle size emulsion.
- (7) RibiTM adjuvant system (RAS), (Ribi Immunochem) containing 2% Squalene, 0.2% Tween 80, and one or more bacterial cell wall components from the group consisting of monophosphoryl lipid A (MPL), trehalose dimycolate (TDM), and cell wall skeleton (CWS), preferably MPL + CWS (DetoxTM); and
- (8) one or more mineral salts (such as an aluminum salt) + a non-toxic derivative of LPS (such as 3dPML).

20 Aluminium salts and MF59 are preferred adjuvants for parenteral immunisation. Mutant bacterial toxins are preferred mucosal adjuvants.

The composition may include an antibiotic.

Further antigens

25 The compositions of the invention may further comprise one or more additional non-GBS antigens, including additional bacterial, viral or parasitic antigens.

In another embodiment, the GBS antigen combinations of the invention are combined with one or more additional, non-GBS antigens suitable for use in a vaccine designed to protect elderly or immunocomprised individuals. For example, the GBS antigen combinations may be combined with an antigen derived from the group consisting of *Enterococcus faecalis*, *Staphylococcus aureus*, 30 *Staphylococcus epidermidis*, *Pseudomonas aeruginosa*, *Legionella pneumophila*, *Listeria monocytogenes*, *Neisseria meningitidis*, influenza, and Parainfluenza virus ('PIV').

Where a saccharide or carbohydrate antigen is used, it is preferably conjugated to a carrier protein in order to enhance immunogenicity {e.g. refs. 42 to 51}. Preferred carrier proteins are bacterial toxins or toxoids, such as diphtheria or tetanus toxoids. The CRM₁₉₇ diphtheria toxoid is 35 particularly preferred {52}. Other carrier polypeptides include the *N.meningitidis* outer membrane

protein {53}, synthetic peptides {54, 55}, heat shock proteins {56, 57}, pertussis proteins {58, 59}, protein D from *H.influenzae* {60}, cytokines {61}, lymphokines, hormones, growth factors, toxin A or B from *C.difficile* {62}, iron-uptake proteins {63}, etc. Where a mixture comprises capsular saccharides from both serogroups A and C, it may be preferred that the ratio (w/w) of MenA 5 saccharide:MenC saccharide is greater than 1 (e.g. 2:1, 3:1, 4:1, 5:1, 10:1 or higher). Different saccharides can be conjugated to the same or different type of carrier protein. Any suitable conjugation reaction can be used, with any suitable linker where necessary.

Toxic protein antigens may be detoxified where necessary e.g. detoxification of pertussis toxin by chemical and/or genetic means.

10 Where a diphtheria antigen is included in the composition it is preferred also to include tetanus antigen and pertussis antigens. Similarly, where a tetanus antigen is included it is preferred also to include diphtheria and pertussis antigens. Similarly, where a pertussis antigen is included it is preferred also to include diphtheria and tetanus antigens.

15 Antigens in the composition will typically be present at a concentration of at least 1 μ g/ml each. In general, the concentration of any given antigen will be sufficient to elicit an immune response against that antigen.

20 As an alternative to using protein antigens in the composition of the invention, nucleic acid encoding the antigen may be used {e.g. refs. 64 to 72}. Protein components of the compositions of the invention may thus be replaced by nucleic acid (preferably DNA e.g. in the form of a plasmid) that encodes the protein.

Definitions

The term "comprising" means "including" as well as "consisting" e.g. a composition "comprising" X may consist exclusively of X or may include something additional e.g. X + Y.

25 The term "about" in relation to a numerical value x means, for example, $x \pm 10\%$.

References to a percentage sequence identity between two amino acid sequences means that, when aligned, that percentage of amino acids are the same in comparing the two sequences. This alignment and the percent homology or sequence identity can be determined using software programs known in the art, for example those described in section 7.7.18 of reference 73. A preferred 30 alignment is determined by the Smith-Waterman homology search algorithm using an affine gap search with a gap open penalty of 12 and a gap extension penalty of 2, BLOSUM matrix of 62. The Smith-Waterman homology search algorithm is disclosed in reference 74.

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